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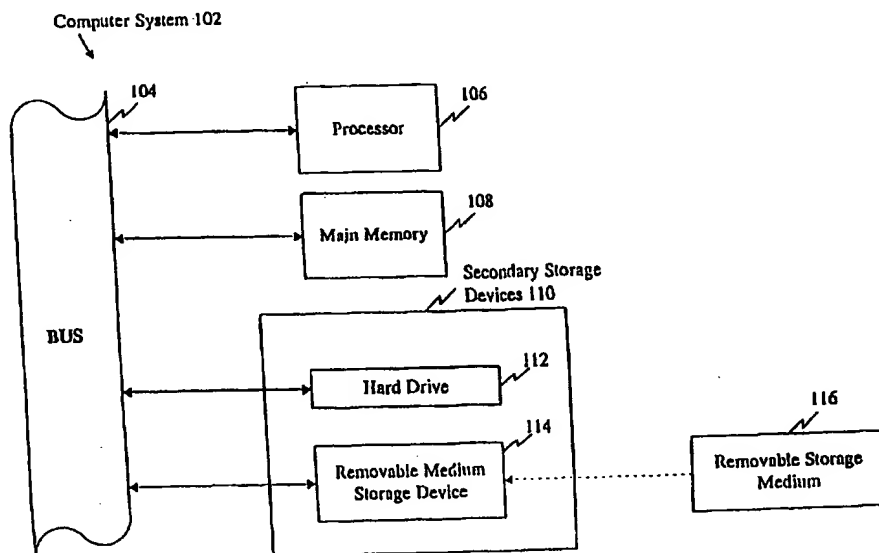
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(54) Title: **STREPTOCOCCUS PNEUMONIAE POLYNUCLEOTIDES AND SEQUENCES**



(57) Abstract

The present invention provides polynucleotide sequences of the genome of *Streptococcus pneumoniae*, polypeptide sequences encoded by the polynucleotide sequences, corresponding polynucleotides and polypeptides, vectors and hosts comprising the polynucleotides, and assays and other uses thereof. The present invention further provides polynucleotide and polypeptide sequence information stored on computer readable media, and computer-based systems and methods which facilitate its use.

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Streptococcus pneumoniae Polynucleotides and Sequences

FIELD OF THE INVENTION

5 The present invention relates to the field of molecular biology. In particular, it relates to, among other things, nucleotide sequences of *Streptococcus pneumoniae*, contigs, ORFs, fragments, probes, primers and related polynucleotides thereof, peptides and polypeptides encoded by the sequences, and uses of the polynucleotides and sequences thereof, such as in fermentation, 10 polypeptide production, assays and pharmaceutical development, among others.

BACKGROUND OF THE INVENTION

15 *Streptococcus pneumoniae* has been one of the most extensively studied microorganisms since its first isolation in 1881. It was the object of many investigations that led to important scientific discoveries. In 1928, Griffith observed that when heat-killed encapsulated pneumococci and live strains constitutiveiy lacking any capsule were concomitantly injected into mice, the nonencapsulated could be converted into encapsulated pneumococci with the same 20 capsular type as the heat-killed strain. Years later, the nature of this "transforming principle," or carrier of genetic information, was shown to be DNA. (Avery, O.T., *et al.*, *J. Exp. Med.*, 79:137-157 (1944)).

25 In spite of the vast number of publications on *S. pneumoniae* many questions about its virulence are still unanswered, and this pathogen remains a major causative agent of serious human disease, especially community-acquired pneumonia. (Johnston, R.B., *et al.*, *Rev. Infect. Dis.* 13(Suppl. 6):S509-517 (1991)). In addition, in developing countries, the pneumococcus is responsible for the death of a large number of children under the age of 5 years from pneumococcal pneumonia. The incidence of pneumococcal disease is highest in infants under 2 30 years of age and in people over 60 years of age. Pneumococci are the second most frequent cause (after *Haemophilus influenzae* type b) of bacterial meningitis and otitis media in children. With the recent introduction of conjugate vaccines for *H. influenzae* type b, pneumococcal meningitis is likely to become increasingly prominent. *S. pneumoniae* is the most important etiologic agent of community-

acquired pneumonia in adults and is the second most common cause of bacterial meningitis behind *Neisseria meningitidis*.

The antibiotic generally prescribed to treat *S. pneumoniae* is benzylpenicillin, although resistance to this and to other antibiotics is found occasionally. Pneumococcal resistance to penicillin results from mutations in its penicillin-binding proteins. In uncomplicated pneumococcal pneumonia caused by a sensitive strain, treatment with penicillin is usually successful unless started too late. Erythromycin or clindamycin can be used to treat pneumonia in patients hypersensitive to penicillin, but resistant strains to these drugs exist. Broad spectrum antibiotics (e.g., the tetracyclines) may also be effective, although tetracycline-resistant strains are not rare. In spite of the availability of antibiotics, the mortality of pneumococcal bacteremia in the last four decades has remained stable between 25 and 29%. (Gillespie, S.H., *et al.*, *J. Med. Microbiol.* 28:237-248 (1989)).

S. pneumoniae is carried in the upper respiratory tract by many healthy individuals. It has been suggested that attachment of pneumococci is mediated by a disaccharide receptor on fibronectin, present on human pharyngeal epithelial cells. (Anderson, B.J., *et al.*, *J. Immunol.* 142:2464-2468 (1989)). The mechanisms by which pneumococci translocate from the nasopharynx to the lung, thereby causing pneumonia, or migrate to the blood, giving rise to bacteremia or septicemia, are poorly understood. (Johnston, R.B., *et al.*, *Rev. Infect. Dis.* 13(Suppl. 6):S509-517 (1991)).

Various proteins have been suggested to be involved in the pathogenicity of *S. pneumoniae*, however, only a few of them have actually been confirmed as virulence factors. Pneumococci produce an IgA1 protease that might interfere with host defense at mucosal surfaces. (Kornfield, S.J., *et al.*, *Rev. Inf. Dis.* 3:521-534 (1981)). *S. pneumoniae* also produces neuraminidase, an enzyme that may facilitate attachment to epithelial cells by cleaving sialic acid from the host glycolipids and gangliosides. Partially purified neuraminidase was observed to induce meningitis-like symptoms in mice; however, the reliability of this finding has been questioned because the neuraminidase preparations used were probably contaminated with cell wall products. Other pneumococcal proteins besides neuraminidase are involved in the adhesion of pneumococci to epithelial and endothelial cells. These pneumococcal proteins have as yet not been identified. Recently, Cundell *et. al.*, reported that peptide permeases can modulate

pneumococcal adherence to epithelial and endothelial cells. It was, however, unclear whether these permeases function directly as adhesions or whether they enhance adherence by modulating the expression of pneumococcal adhesions. (DeVelasco, E.A., *et al.*, *Micro. Rev.* 59:591-603 (1995). A better understanding 5 of the virulence factors determining its pathogenicity will need to be developed to cope with the devastating effects of pneumococcal disease in humans.

Ironically, despite the prominent role of *S. pneumoniae* in the discovery of DNA, little is known about the molecular genetics of the organism. The *S. pneumoniae* genome consists of one circular, covalently closed, double-stranded 10 DNA and a collection of so-called variable accessory elements, such as prophages, plasmids, transposons and the like. Most physical characteristics and almost all of the genes of *S. pneumoniae* are unknown. Among the few that have been identified, most have not been physically mapped or characterized in detail. Only a few genes of this organism have been sequenced. (See, for instance current 15 versions of GENBANK and other nucleic acid databases, and references that relate to the genome of *S. pneumoniae* such as those set out elsewhere herein.)

It is clear that the etiology of diseases mediated or exacerbated by *S. pneumoniae*, infection involves the programmed expression of *S. pneumoniae* genes, and that characterizing the genes and their patterns of expression would add 20 dramatically to our understanding of the organism and its host interactions. Knowledge of *S. pneumoniae* genes and genomic organization would improve our understanding of disease etiology and lead to improved and new ways of preventing, ameliorating, arresting and reversing diseases. Moreover, characterized genes and genomic fragments of *S. pneumoniae* would provide 25 reagents for, among other things, detecting, characterizing and controlling *S. pneumoniae* infections. There is a need to characterize the genome of *S. pneumoniae* and for polynucleotides of this organism.

SUMMARY OF THE INVENTION

5 The present invention is based on the sequencing of fragments of the *Streptococcus pneumoniae* genome. The primary nucleotide sequences which were generated are provided in SEQ ID NOS:1-391.

10 The present invention provides the nucleotide sequence of several hundred contigs of the *Streptococcus pneumoniae* genome, which are listed in tables below and set out in the Sequence Listing submitted herewith, and representative fragments thereof, in a form which can be readily used, analyzed, and interpreted by a skilled artisan. In one embodiment, the present invention is provided as contiguous strings of primary sequence information corresponding to the nucleotide sequences depicted in SEQ ID NOS:1-391.

15 The present invention further provides nucleotide sequences which are at least 95% identical to the nucleotide sequences of SEQ ID NOS:1-391.

20 The nucleotide sequence of SEQ ID NOS:1-391, a representative fragment thereof, or a nucleotide sequence which is at least 95% identical to the nucleotide sequence of SEQ ID NOS:1-391 may be provided in a variety of mediums to facilitate its use. In one application of this embodiment, the sequences of the present invention are recorded on computer readable media. Such media includes, but is not limited to: magnetic storage media, such as floppy discs, hard disc storage medium, and magnetic tape; optical storage media such as CD-ROM; electrical storage media such as RAM and ROM; and hybrids of these categories such as magnetic/optical storage media.

25 The present invention further provides systems, particularly computer-based systems which contain the sequence information herein described stored in a data storage means. Such systems are designed to identify commercially important fragments of the *Streptococcus pneumoniae* genome.

30 Another embodiment of the present invention is directed to fragments of the *Streptococcus pneumoniae* genome having particular structural or functional attributes. Such fragments of the *Streptococcus pneumoniae* genome of the present invention include, but are not limited to, fragments which encode peptides, hereinafter referred to as open reading frames or ORFs, fragments which modulate the expression of an operably linked ORF, hereinafter referred to as expression modulating fragments or EMFs, and fragments which can be used to diagnose the

presence of *Streptococcus pneumoniae* in a sample, hereinafter referred to as diagnostic fragments or DFs.

Each of the ORFs in fragments of the *Streptococcus pneumoniae* genome disclosed in Tables 1-3, and the EMFs found 5' to the ORFs, can be used in numerous ways as polynucleotide reagents. For instance, the sequences can be used as diagnostic probes or amplification primers for detecting or determining the presence of a specific microbe in a sample, to selectively control gene expression in a host and in the production of polypeptides, such as polypeptides encoded by ORFs of the present invention, particular those polypeptides that have a pharmacological activity.

The present invention further includes recombinant constructs comprising one or more fragments of the *Streptococcus pneumoniae* genome of the present invention. The recombinant constructs of the present invention comprise vectors, such as a plasmid or viral vector, into which a fragment of the *Streptococcus pneumoniae* has been inserted.

The present invention further provides host cells containing any of the isolated fragments of the *Streptococcus pneumoniae* genome of the present invention. The host cells can be a higher eukaryotic host cell, such as a mammalian cell, a lower eukaryotic cell, such as a yeast cell, or a prokaryotic cell such as a bacterial cell.

The present invention is further directed to isolated polypeptides and proteins encoded by ORFs of the present invention. A variety of methods, well known to those of skill in the art, routinely may be utilized to obtain any of the polypeptides and proteins of the present invention. For instance, polypeptides and proteins of the present invention having relatively short, simple amino acid sequences readily can be synthesized using commercially available automated peptide synthesizers. Polypeptides and proteins of the present invention also may be purified from bacterial cells which naturally produce the protein. Yet another alternative is to purify polypeptide and proteins of the present invention from cells which have been altered to express them.

The invention further provides methods of obtaining homologs of the fragments of the *Streptococcus pneumoniae* genome of the present invention and homologs of the proteins encoded by the ORFs of the present invention. Specifically, by using the nucleotide and amino acid sequences disclosed herein as

a probe or as primers, and techniques such as PCR cloning and colony/plaque hybridization, one skilled in the art can obtain homologs.

The invention further provides antibodies which selectively bind polypeptides and proteins of the present invention. Such antibodies include both 5 monoclonal and polyclonal antibodies.

The invention further provides hybridomas which produce the above-described antibodies. A hybridoma is an immortalized cell line which is capable of secreting a specific monoclonal antibody.

The present invention further provides methods of identifying test samples 10 derived from cells which express one of the ORFs of the present invention, or a homolog thereof. Such methods comprise incubating a test sample with one or more of the antibodies of the present invention, or one or more of the DFs of the present invention, under conditions which allow a skilled artisan to determine if the sample contains the ORF or product produced therefrom.

15 In another embodiment of the present invention, kits are provided which contain the necessary reagents to carry out the above-described assays.

Specifically, the invention provides a compartmentalized kit to receive, in close confinement, one or more containers which comprises: (a) a first container 20 comprising one of the antibodies, or one of the DFs of the present invention; and (b) one or more other containers comprising one or more of the following: wash reagents, reagents capable of detecting presence of bound antibodies or hybridized DFs.

Using the isolated proteins of the present invention, the present invention further provides methods of obtaining and identifying agents capable of binding to 25 a polypeptide or protein encoded by one of the ORFs of the present invention. Specifically, such agents include, as further described below, antibodies, peptides, carbohydrates, pharmaceutical agents and the like. Such methods comprise steps of: (a) contacting an agent with an isolated protein encoded by one of the ORFs of the present invention; and (b) determining whether the agent binds to said protein.

30 The present genomic sequences of *Streptococcus pneumoniae* will be of great value to all laboratories working with this organism and for a variety of commercial purposes. Many fragments of the *Streptococcus pneumoniae* genome will be immediately identified by similarity searches against GenBank or protein databases and will be of immediate value to *Streptococcus pneumoniae* researchers

and for immediate commercial value for the production of proteins or to control gene expression.

The methodology and technology for elucidating extensive genomic sequences of bacterial and other genomes has and will greatly enhance the ability to 5 analyze and understand chromosomal organization. In particular, sequenced contigs and genomes will provide the models for developing tools for the analysis of chromosome structure and function, including the ability to identify genes within large segments of genomic DNA, the structure, position, and spacing of regulatory elements, the identification of genes with potential industrial applications, and the 10 ability to do comparative genomic and molecular phylogeny.

DESCRIPTION OF THE FIGURES

15 **FIGURE 1** is a block diagram of a computer system (102) that can be used to implement computer-based systems of present invention.

20 **FIGURE 2** is a schematic diagram depicting the data flow and computer programs used to collect, assemble, edit and annotate the contigs of the *Streptococcus pneumoniae* genome of the present invention. Both Macintosh and Unix platforms are used to handle the AB 373 and 377 sequence data files, largely as described in Kerlavage *et al.*, *Proceedings of the Twenty-Sixth Annual Hawaii International Conference on System Sciences*, 585, IEEE Computer Society Press, Washington D.C. (1993). Factura (AB) is a Macintosh program designed for 25 automatic vector sequence removal and end-trimming of sequence files. The program Loadis runs on a Macintosh platform and parses the feature data extracted from the sequence files by Factura to the Unix based *Streptococcus pneumoniae* relational database. Assembly of contigs (and whole genome sequences) is accomplished by retrieving a specific set of sequence files and their associated features using Extrseq, a Unix utility for retrieving sequences from an SQL 30 database. The resulting sequence file is processed by seq_filter to trim portions of the sequences with more than 2% ambiguous nucleotides. The sequence files were assembled using TIGR Assembler, an assembly engine designed at The Institute for Genomic Research (TIGR) for rapid and accurate assembly of thousands of sequence fragments. The collection of contigs generated by the assembly step is 35 loaded into the database with the lassie program. Identification of open reading

frames (ORFs) is accomplished by processing contigs with zorf or GenMark. The ORFs are searched against *S. pneumoniae* sequences from GenBank and against all protein sequences using the BLASTN and BLASTP programs, described in Altschul *et al.*, *J. Mol. Biol.* 215: 403-410 (1990)). Results of the ORF 5 determination and similarity searching steps were loaded into the database. As described below, some results of the determination and the searches are set out in Tables 1-3.

DETAILED DESCRIPTION OF ILLUSTRATIVE EMBODIMENTS

10

The present invention is based on the sequencing of fragments of the *Streptococcus pneumoniae* genome and analysis of the sequences. The primary nucleotide sequences generated by sequencing the fragments are provided in SEQ ID NOS:1-391. (As used herein, the "primary sequence" refers to the nucleotide 15 sequence represented by the IUPAC nomenclature system.)

In addition to the aforementioned *Streptococcus pneumoniae* polynucleotide and polynucleotide sequences, the present invention provides the nucleotide sequences of SEQ ID NOS:1-391, or representative fragments thereof, in a form which can be readily used, analyzed, and interpreted by a skilled artisan.

20

As used herein, a "representative fragment of the nucleotide sequence depicted in SEQ ID NOS:1-391" refers to any portion of the SEQ ID NOS:1-391 which is not presently represented within a publicly available database. Preferred representative fragments of the present invention are *Streptococcus pneumoniae* open reading frames (ORFs), expression modulating fragment (EMFs) and 25 fragments which can be used to diagnose the presence of *Streptococcus pneumoniae* in sample (DFs). A non-limiting identification of preferred representative fragments is provided in Tables 1-3. As discussed in detail below, the information provided in SEQ ID NOS:1-391 and in Tables 1-3 together with routine cloning, synthesis, sequencing and assay methods will enable those skilled 30 in the art to clone and sequence all "representative fragments" of interest, including open reading frames encoding a large variety of *Streptococcus pneumoniae* proteins.

While the presently disclosed sequences of SEQ ID NOS:1-391 are highly accurate, sequencing techniques are not perfect and, in relatively rare instances, 35 further investigation of a fragment or sequence of the invention may reveal a

nucleotide sequence error present in a nucleotide sequence disclosed in SEQ ID NOS:1-391. However, once the present invention is made available (*i.e.*, once the information in SEQ ID NOS:1-391 and Tables 1-3 has been made available), resolving a rare sequencing error in SEQ ID NOS:1-391 will be well within the 5 skill of the art. The present disclosure makes available sufficient sequence information to allow any of the described contigs or portions thereof to be obtained readily by straightforward application of routine techniques. Further sequencing of such polynucleotide may proceed in like manner using manual and automated sequencing methods which are employed ubiquitous in the art. Nucleotide 10 sequence editing software is publicly available. For example, Applied Biosystem's (AB) AutoAssembler can be used as an aid during visual inspection of nucleotide sequences. By employing such routine techniques potential errors readily may be identified and the correct sequence then may be ascertained by targeting further sequencing effort, also of a routine nature, to the region containing the potential 15 error.

Even if all of the very rare sequencing errors in SEQ ID NOS:1-391 were corrected, the resulting nucleotide sequences would still be at least 95% identical, nearly all would be at least 99% identical, and the great majority would be at least 99.9% identical to the nucleotide sequences of SEQ ID NOS:1-391.

As discussed elsewhere herein, polynucleotides of the present invention readily may be obtained by routine application of well known and standard procedures for cloning and sequencing DNA. Detailed methods for obtaining 20 libraries and for sequencing are provided below, for instance. A wide variety of *Streptococcus pneumoniae* strains that can be used to prepare *S. pneumoniae* 25 genomic DNA for cloning and for obtaining polynucleotides of the present invention are available to the public from recognized depository institutions, such as the American Type Culture Collection (ATCC). While the present invention is enabled by the sequences and other information herein disclosed, the *S. pneumoniae* strain that provided the DNA of the present Sequence Listing, Strain 30 7/87 14.8.91, has been deposited in the ATCC, as a convenience to those of skill in the art. As a further convenience, a library of *S. pneumoniae* genomic DNA, derived from the same strain, also has been deposited in the ATCC. The *S. pneumoniae* strain was deposited on October 10, 1996, and was given Deposit No. 35 55840, and the cDNA library was deposited on October 11, 1996 and was given Deposit No. 97755. The genomic fragments in the library are 15 to 20 kb

fragments generated by partial Sau3A1 digestion and they are inserted into the BamHI site in the well-known lambda-derived vector lambda DASH II (Stratagene, La Jolla, CA). The provision of the deposits is not a waiver of any rights of the inventors or their assignees in the present subject matter.

5 The nucleotide sequences of the genomes from different strains of *Streptococcus pneumoniae* differ somewhat. However, the nucleotide sequences of the genomes of all *Streptococcus pneumoniae* strains will be at least 95% identical, in corresponding part, to the nucleotide sequences provided in SEQ ID NOS:1-391. Nearly all will be at least 99% identical and the great majority will be
10 99.9% identical.

Thus, the present invention further provides nucleotide sequences which are at least 95%, preferably 99% and most preferably 99.9% identical to the nucleotide sequences of SEQ ID NOS:1-391, in a form which can be readily used, analyzed and interpreted by the skilled artisan.

15 Methods for determining whether a nucleotide sequence is at least 95%, at least 99% or at least 99.9% identical to the nucleotide sequences of SEQ ID NOS:1-391 are routine and readily available to the skilled artisan. For example, the well known fasta algorithm described in Pearson and Lipman, *Proc. Natl. Acad. Sci. USA* 85: 2444 (1988) can be used to generate the percent identity of nucleotide
20 sequences. The BLASTN program also can be used to generate an identity score of polynucleotides compared to one another.

COMPUTER RELATED EMBODIMENTS

25 The nucleotide sequences provided in SEQ ID NOS:1-391, a representative fragment thereof, or a nucleotide sequence at least 95%, preferably at least 99% and most preferably at least 99.9% identical to a polynucleotide sequence of SEQ ID NOS:1-391 may be "provided" in a variety of mediums to facilitate use thereof. As used herein, provided refers to a manufacture, other than an isolated nucleic acid molecule, which contains a nucleotide sequence of the present invention; *i.e.*,
30 a nucleotide sequence provided in SEQ ID NOS:1-391, a representative fragment thereof, or a nucleotide sequence at least 95%, preferably at least 99% and most preferably at least 99.9% identical to a polynucleotide of SEQ ID NOS:1-391. Such a manufacture provides a large portion of the *Streptococcus pneumoniae* genome and parts thereof (*e.g.*, a *Streptococcus pneumoniae* open reading frame
35 (ORF)) in a form which allows a skilled artisan to examine the manufacture using

means not directly applicable to examining the *Streptococcus pneumoniae* genome or a subset thereof as it exists in nature or in purified form.

In one application of this embodiment, a nucleotide sequence of the present invention can be recorded on computer readable media. As used herein, "computer readable media" refers to any medium which can be read and accessed directly by a computer. Such media include, but are not limited to: magnetic storage media, such as floppy discs, hard disc storage medium, and magnetic tape; optical storage media such as CD- ROM; electrical storage media such as RAM and ROM; and hybrids of these categories, such as magnetic/optical storage media. A skilled artisan can readily appreciate how any of the presently known computer readable mediums can be used to create a manufacture comprising computer readable medium having recorded thereon a nucleotide sequence of the present invention. Likewise, it will be clear to those of skill how additional computer readable media that may be developed also can be used to create analogous manufactures having recorded thereon a nucleotide sequence of the present invention.

As used herein, "recorded" refers to a process for storing information on computer readable medium. A skilled artisan can readily adopt any of the presently known methods for recording information on computer readable medium to generate manufactures comprising the nucleotide sequence information of the present invention. A variety of data storage structures are available to a skilled artisan for creating a computer readable medium having recorded thereon a nucleotide sequence of the present invention. The choice of the data storage structure will generally be based on the means chosen to access the stored information. In addition, a variety of data processor programs and formats can be used to store the nucleotide sequence information of the present invention on computer readable medium. The sequence information can be represented in a word processing text file, formatted in commercially- available software such as WordPerfect and MicroSoft Word, or represented in the form of an ASCII file, stored in a database application, such as DB2, Sybase, Oracle, or the like. A skilled artisan can readily adapt any number of data-processor structuring formats (e.g., text file or database) in order to obtain computer readable medium having recorded thereon the nucleotide sequence information of the present invention.

Computer software is publicly available which allows a skilled artisan to access sequence information provided in a computer readable medium. Thus, by providing in computer readable form the nucleotide sequences of SEQ ID NOS:1-

391, a representative fragment thereof, or a nucleotide sequence at least 95%, preferably at least 99% and most preferably at least 99.9% identical to a sequence of SEQ ID NOS:1-391 the present invention enables the skilled artisan routinely to access the provided sequence information for a wide variety of purposes.

5 The examples which follow demonstrate how software which implements the BLAST (Altschul *et al.*, *J. Mol. Biol.* 215:403-410 (1990)) and BLAZE (Brutlag *et al.*, *Comp. Chem.* 17:203-207 (1993)) search algorithms on a Sybase system was used to identify open reading frames (ORFs) within the *Streptococcus pneumoniae* genome which contain homology to ORFs or proteins from both
10 *Streptococcus pneumoniae* and from other organisms. Among the ORFs discussed herein are protein encoding fragments of the *Streptococcus pneumoniae* genome useful in producing commercially important proteins, such as enzymes used in fermentation reactions and in the production of commercially useful metabolites.

15 The present invention further provides systems, particularly computer-based systems, which contain the sequence information described herein. Such systems are designed to identify, among other things, commercially important fragments of the *Streptococcus pneumoniae* genome.

20 As used herein, "a computer-based system" refers to the hardware means, software means, and data storage means used to analyze the nucleotide sequence information of the present invention. The minimum hardware means of the computer-based systems of the present invention comprises a central processing unit (CPU), input means, output means, and data storage means. A skilled artisan can readily appreciate that any one of the currently available computer-based systems are suitable for use in the present invention.

25 As stated above, the computer-based systems of the present invention comprise a data storage means having stored therein a nucleotide sequence of the present invention and the necessary hardware means and software means for supporting and implementing a search means.

30 As used herein, "data storage means" refers to memory which can store nucleotide sequence information of the present invention, or a memory access means which can access manufactures having recorded thereon the nucleotide sequence information of the present invention.

35 As used herein, "search means" refers to one or more programs which are implemented on the computer-based system to compare a target sequence or target structural motif with the sequence information stored within the data storage

means. Search means are used to identify fragments or regions of the present genomic sequences which match a particular target sequence or target motif. A variety of known algorithms are disclosed publicly and a variety of commercially available software for conducting search means are and can be used in the 5 computer-based systems of the present invention. Examples of such software includes, but is not limited to, MacPattern (EMBL), BLASTN and BLASTX (NCBIA). A skilled artisan can readily recognize that any one of the available algorithms or implementing software packages for conducting homology searches can be adapted for use in the present computer-based systems.

10 As used herein, a "target sequence" can be any DNA or amino acid sequence of six or more nucleotides or two or more amino acids. A skilled artisan can readily recognize that the longer a target sequence is, the less likely a target sequence will be present as a random occurrence in the database. The most preferred sequence length of a target sequence is from about 10 to 100 amino acids 15 or from about 30 to 300 nucleotide residues. However, it is well recognized that searches for commercially important fragments, such as sequence fragments involved in gene expression and protein processing, may be of shorter length.

20 As used herein, "a target structural motif," or "target motif," refers to any rationally selected sequence or combination of sequences in which the sequence(s) 25 are chosen based on a three-dimensional configuration which is formed upon the folding of the target motif. There are a variety of target motifs known in the art. Protein target motifs include, but are not limited to, enzymic active sites and signal sequences. Nucleic acid target motifs include, but are not limited to, promoter sequences, hairpin structures and inducible expression elements (protein binding sequences).

30 A variety of structural formats for the input and output means can be used to input and output the information in the computer-based systems of the present invention. A preferred format for an output means ranks fragments of the *Streptococcus pneumoniae* genomic sequences possessing varying degrees of 35 homology to the target sequence or target motif. Such presentation provides a skilled artisan with a ranking of sequences which contain various amounts of the target sequence or target motif and identifies the degree of homology contained in the identified fragment.

35 A variety of comparing means can be used to compare a target sequence or target motif with the data storage means to identify sequence fragments of the

5 *Streptococcus pneumoniae* genome. In the present examples, implementing software which implement the BLAST and BLAZE algorithms, described in Altschul *et al.*, *J. Mol. Biol.* 215: 403-410 (1990), is used to identify open reading frames within the *Streptococcus pneumoniae* genome. A skilled artisan can readily recognize that any one of the publicly available homology search programs can be used as the search means for the computer-based systems of the present invention. Of course, suitable proprietary systems that may be known to those of skill also may be employed in this regard.

10 Figure 1 provides a block diagram of a computer system illustrative of embodiments of this aspect of present invention. The computer system 102 includes a processor 106 connected to a bus 104. Also connected to the bus 104 are a main memory 108 (preferably implemented as random access memory, RAM) and a variety of secondary storage devices 110, such as a hard drive 112 and a removable medium storage device 114. The removable medium storage device 114 15 may represent, for example, a floppy disk drive, a CD-ROM drive, a magnetic tape drive, *etc.* A removable storage medium 116 (such as a floppy disk, a compact disk, a magnetic tape, *etc.*) containing control logic and/or data recorded therein may be inserted into the removable medium storage device 114. The computer system 102 includes appropriate software for reading the control logic and/or the 20 data from the removable medium storage device 114, once it is inserted into the removable medium storage device 114.

25 A nucleotide sequence of the present invention may be stored in a well known manner in the main memory 108, any of the secondary storage devices 110, and/or a removable storage medium 116. During execution, software for accessing and processing the genomic sequence (such as search tools, comparing tools, *etc.*) reside in main memory 108, in accordance with the requirements and operating parameters of the operating system, the hardware system and the software program or programs.

BIOCHEMICAL EMBODIMENTS

Other embodiments of the present invention are directed to isolated fragments of the *Streptococcus pneumoniae* genome. The fragments of the 5 *Streptococcus pneumoniae* genome of the present invention include, but are not limited to fragments which encode peptides and polypeptides, hereinafter open reading frames (ORFs), fragments which modulate the expression of an operably linked ORF, hereinafter expression modulating fragments (EMFs) and fragments which can be used to diagnose the presence of *Streptococcus pneumoniae* in a 10 sample, hereinafter diagnostic fragments (DFs).

As used herein, an "isolated nucleic acid molecule" or an "isolated fragment 15 of the *Streptococcus pneumoniae* genome" refers to a nucleic acid molecule possessing a specific nucleotide sequence which has been subjected to purification means to reduce, from the composition, the number of compounds which are normally associated with the composition. Particularly, the term refers to the nucleic acid molecules having the sequences set out in SEQ ID NOS:1-391, to representative fragments thereof as described above, to polynucleotides at least 95%, preferably at least 99% and especially preferably at least 99.9% identical in sequence thereto, also as set out above.

20 A variety of purification means can be used to generate the isolated fragments of the present invention. These include, but are not limited to methods which separate constituents of a solution based on charge, solubility, or size.

In one embodiment, *Streptococcus pneumoniae* DNA can be enzymatically 25 sheared to produce fragments of 15-20 kb in length. These fragments can then be used to generate a *Streptococcus pneumoniae* library by inserting them into lambda clones as described in the Examples below. Primers flanking, for example, an ORF, such as those enumerated in Tables 1-3 can then be generated using nucleotide sequence information provided in SEQ ID NOS:1-391. Well known and routine techniques of PCR cloning then can be used to isolate the ORF from 30 the lambda DNA library or *Streptococcus pneumoniae* genomic DNA. Thus, given the availability of SEQ ID NOS:1-391, the information in Tables 1, 2 and 3, and the information that may be obtained readily by analysis of the sequences of SEQ ID NOS:1-391 using methods set out above, those of skill will be enabled by the 35 present disclosure to isolate any ORF-containing or other nucleic acid fragment of the present invention.

The isolated nucleic acid molecules of the present invention include, but are not limited to single stranded and double stranded DNA, and single stranded RNA.

As used herein, an "open reading frame," ORF, means a series of triplets coding for amino acids without any termination codons and is a sequence 5 translatable into protein.

Tables 1, 2, and 3 list ORFs in the *Streptococcus pneumoniae* genomic contigs of the present invention that were identified as putative coding regions by the GeneMark software using organism-specific second-order Markov probability transition matrices. It will be appreciated that other criteria can be used, in 10 accordance with well known analytical methods, such as those discussed herein, to generate more inclusive, more restrictive, or more selective lists.

Table 1 sets out ORFs in the *Streptococcus pneumoniae* contigs of the present invention that over a continuous region of at least 50 bases are 95% or more identical (by BLAST analysis) to a nucleotide sequence available through 15 GenBank in October, 1997.

Table 2 sets out ORFs in the *Streptococcus pneumoniae* contigs of the present invention that are not in Table 1 and match, with a BLASTP probability score of 0.01 or less, a polypeptide sequence available through GenBank in October, 1997.

Table 3 sets out ORFs in the *Streptococcus pneumoniae* contigs of the present invention that do not match significantly, by BLASTP analysis, a polypeptide sequence available through GenBank in October, 1997.

In each table, the first and second columns identify the ORF by, respectively, contig number and ORF number within the contig; the third column 25 indicates the first nucleotide of the ORF (actually the first nucleotide of the stop codon immediately preceding the ORF), counting from the 5' end of the contig strand; and the fourth column, "stop (nt)" indicates the last nucleotide of the stop codon defining the 3' end of the ORF.

In Tables 1 and 2, column five, lists the Reference for the closest 30 matching sequence available through GenBank. These reference numbers are the databases entry numbers commonly used by those of skill in the art, who will be familiar with their denominators. Descriptions of the nomenclature are available from the National Center for Biotechnology Information. Column six in Tables 1 and 2 provides the gene name of the matching sequence; column seven provides 35 the BLAST identity score and column eight the BLAST similarity score from the

comparison of the ORF and the homologous gene; and column nine indicates the length in nucleotides of the highest scoring segment pair identified by the BLAST identity analysis.

Each ORF described in the tables is defined by "start (nt)" (5') and "stop (nt)" (3') nucleotide position numbers. These position numbers refer to the boundaries of each ORF and provide orientation with respect to whether the forward or reverse strand is the coding strand and which reading frame the coding sequence is contained. The "start" position is the first nucleotide of the triplet encoding a stop codon just 5' to the ORF and the "stop" position is the last 10 nucleotide of the triplet encoding the next in-frame stop codon (i.e., the stop codon at the 3' end of the ORF). Those of ordinary skill in the art appreciate that preferred fragments within each ORF described in the table include fragments of each ORF which include the entire sequence from the delineated "start" and "stop" positions excepting the first and last three nucleotides since these encode stop 15 codons. Thus, polynucleotides set out as ORFs in the tables but lacking the three (3) 5' nucleotides and the three (3) 3' nucleotides are encompassed by the present invention. Those of skill also appreciate that particularly preferred are fragments within each ORF that are polynucleotide fragments comprising polypeptide coding sequence. As defined herein, "coding sequence" includes the fragment within an 20 ORF beginning at the first in-frame ATG (triplet encoding methionine) and ending with the last nucleotide prior to the triplet encoding the 3' stop codon. Preferred are fragments comprising the entire coding sequence and fragments comprising the entire coding sequence, excepting the coding sequence for the N-terminal methionine. Those of skill appreciate that the N-terminal methionine is often 25 removed during post-translational processing and that polynucleotides lacking the ATG can be used to facilitate production of N-terminal fusion proteins which may be beneficial in the production or use of genetically engineered proteins. Of course, due to the degeneracy of the genetic code many polynucleotides can encode a given polypeptide. Thus, the invention further includes polynucleotides comprising a 30 nucleotide sequence encoding a polypeptide sequence itself encoded by the coding sequence within an ORF described in Tables 1-3 herein. Further, polynucleotides at least 95%, preferably at least 99% and especially preferably at least 99.9% identical in sequence to the foregoing polynucleotides, are contemplated by the present invention.

5 Polypeptides encoded by polynucleotides described above and elsewhere herein are also provided by the present invention as are polypeptide comprising a an amino acid sequence at least about 95%, preferably at least 97% and even more preferably 99% identical to the amino acid sequence of a polypeptide encoded by an
ORF shown in Tables 1-3. These polypeptides may or may not comprise an N-terminal methionine.

10 The concepts of percent identity and percent similarity of two polypeptide sequences is well understood in the art. For example, two polypeptides 10 amino acids in length which differ at three amino acid positions (e.g., at positions 1, 3 and 5) are said to have a percent identity of 70%. However, the same two polypeptides would be deemed to have a percent similarity of 80% if, for example at position 5, the amino acids moieties, although not identical, were "similar" (i.e., possessed similar biochemical characteristics). Many programs for analysis of nucleotide or amino acid sequence similarity, such as fasta and BLAST specifically
15 list percent identity of a matching region as an output parameter. Thus, for instance, Tables 1 and 2 herein enumerate the percent identity of the highest scoring segment pair in each ORF and its listed relative. Further details concerning the algorithms and criteria used for homology searches are provided below and are described in the pertinent literature highlighted by the citations
20 provided below.

25 It will be appreciated that other criteria can be used to generate more inclusive and more exclusive listings of the types set out in the tables. As those of skill will appreciate, narrow and broad searches both are useful. Thus, a skilled artisan can readily identify ORFs in contigs of the *Streptococcus pneumoniae* genome other than those listed in Tables 1-3, such as ORFs which are overlapping or encoded by the opposite strand of an identified ORF in addition to those ascertainable using the computer-based systems of the present invention.

30 As used herein, an "expression modulating fragment," EMF, means a series of nucleotide molecules which modulates the expression of an operably linked ORF or EMF.

As used herein, a sequence is said to "modulate the expression of an operably linked sequence" when the expression of the sequence is altered by the presence of the EMF. EMFs include, but are not limited to, promoters, and promoter modulating sequences (inducible elements). One class of EMFs are 5 fragments which induce the expression of an operably linked ORF in response to a specific regulatory factor or physiological event.

EMF sequences can be identified within the contigs of the *Streptococcus pneumoniae* genome by their proximity to the ORFs provided in Tables 1-3. An intergenic segment, or a fragment of the intergenic segment, from about 10 to 200 10 nucleotides in length, taken from any one of the ORFs of Tables 1-3 will modulate the expression of an operably linked ORF in a fashion similar to that found with the naturally linked ORF sequence. As used herein, an "intergenic segment" refers to fragments of the *Streptococcus pneumoniae* genome which are between two ORF(s) herein described. EMFs also can be identified using known EMFs as a 15 target sequence or target motif in the computer-based systems of the present invention. Further, the two methods can be combined and used together.

The presence and activity of an EMF can be confirmed using an EMF trap vector. An EMF trap vector contains a cloning site linked to a marker sequence. A 20 marker sequence encodes an identifiable phenotype, such as antibiotic resistance or a complementing nutrition auxotrophic factor, which can be identified or assayed when the EMF trap vector is placed within an appropriate host under appropriate conditions. As described above, a EMF will modulate the expression of an operably linked marker sequence. A more detailed discussion of various marker sequences is provided below. A sequence which is suspected as being an EMF is 25 cloned in all three reading frames in one or more restriction sites upstream from the marker sequence in the EMF trap vector. The vector is then transformed into an appropriate host using known procedures and the phenotype of the transformed host is examined under appropriate conditions. As described above, an EMF will modulate the expression of an operably linked marker sequence.

30 As used herein, a "diagnostic fragment," DF, means a series of nucleotide molecules which selectively hybridize to *Streptococcus pneumoniae* sequences. DFs can be readily identified by identifying unique sequences within contigs of the *Streptococcus pneumoniae* genome, such as by using well-known computer analysis software, and by generating and testing probes or amplification primers

consisting of the DF sequence in an appropriate diagnostic format which determines amplification or hybridization selectivity.

The sequences falling within the scope of the present invention are not limited to the specific sequences herein described, but also include allelic and species variations thereof. Allelic and species variations can be routinely determined by comparing the sequences provided in SEQ ID NOS:1-391, a representative fragment thereof, or a nucleotide sequence at least 95%, preferably at least 99% and most at least preferably 99.9% identical to SEQ ID NOS:1-391, with a sequence from another isolate of the same species. Furthermore, to accommodate codon variability, the invention includes nucleic acid molecules coding for the same amino acid sequences as do the specific ORFs disclosed herein. In other words, in the coding region of an ORF, substitution of one codon for another which encodes the same amino acid is expressly contemplated. Any specific sequence disclosed herein can be readily screened for errors by resequencing a particular fragment, such as an ORF, in both directions (*i.e.*, sequence both strands). Alternatively, error screening can be performed by sequencing corresponding polynucleotides of *Streptococcus pneumoniae* origin isolated by using part or all of the fragments in question as a probe or primer.

Preferred DFs of the present invention comprise at least about 17, preferably at least about 20, and more preferably at least about 50 contiguous nucleotides within an ORF set out in Tables 1-3. Most highly preferred DFs specifically hybridize to a polynucleotide containing the sequence of the ORF from which they are derived. Specific hybridization occurs even under stringent conditions defined elsewhere herein.

Each of the ORFs of the *Streptococcus pneumoniae* genome disclosed in Tables 1, 2 and 3, and the EMFs found 5' to the ORFs, can be used as polynucleotide reagents in numerous ways. For example, the sequences can be used as diagnostic probes or diagnostic amplification primers to detect the presence of a specific microbe in a sample, particularly *Streptococcus pneumoniae*. Especially preferred in this regard are ORFs such as those of Table 3, which do not match previously characterized sequences from other organisms and thus are most likely to be highly selective for *Streptococcus pneumoniae*. Also particularly preferred are ORFs that can be used to distinguish between strains of *Streptococcus pneumoniae*, particularly those that distinguish medically important strain, such as drug-resistant strains.

In addition, the fragments of the present invention, as broadly described, can be used to control gene expression through triple helix formation or antisense DNA or RNA, both of which methods are based on the binding of a polynucleotide sequence to DNA or RNA. Triple helix-formation optimally results in a shut-off of 5 RNA transcription from DNA, while antisense RNA hybridization blocks translation of an mRNA molecule into polypeptide. Information from the sequences of the present invention can be used to design antisense and triple helix-forming oligonucleotides. Polynucleotides suitable for use in these methods are usually 20 to 40 bases in length and are designed to be complementary to a region 10 of the gene involved in transcription, for triple-helix formation, or to the mRNA itself, for antisense inhibition. Both techniques have been demonstrated to be effective in model systems, and the requisite techniques are well known and involve routine procedures. Triple helix techniques are discussed in, for example, Lee *et al.*, *Nucl. Acids Res.* 6:3073 (1979); Cooney *et al.*, *Science* 241:456 15 (1988); and Dervan *et al.*, *Science* 251:1360 (1991). Antisense techniques in general are discussed in, for instance, Okano, *J. Neurochem.* 56:560 (1991) and *Oligodeoxynucleotides as Antisense Inhibitors of Gene Expression*, CRC Press, Boca Raton, FL (1988)).

The present invention further provides recombinant constructs comprising 20 one or more fragments of the *Streptococcus pneumoniae* genomic fragments and contigs of the present invention. Certain preferred recombinant constructs of the present invention comprise a vector, such as a plasmid or viral vector, into which a fragment of the *Streptococcus pneumoniae* genome has been inserted, in a forward or reverse orientation. In the case of a vector comprising one of the ORFs of the 25 present invention, the vector may further comprise regulatory sequences, including for example, a promoter, operably linked to the ORF. For vectors comprising the EMFs of the present invention, the vector may further comprise a marker sequence or heterologous ORF operably linked to the EMF.

Large numbers of suitable vectors and promoters are known to those of 30 skill in the art and are commercially available for generating the recombinant constructs of the present invention. The following vectors are provided by way of example. Useful bacterial vectors include phagescript, PsiX174, pBluescript SK, pBS KS, pNH8a, pNH16a, pNH18a, pNH46a (available from Stratagene); pTrc99A, pKK223-3, pKK233-3, pDR540, pRIT5 (available from Pharmacia). 35 Useful eukaryotic vectors include pWLneo, pSV2cat, pOG44, pXT1, pSG

(available from Stratagene) pSVK3, pBPV, pMSG, pSVL (available from Pharmacia).

Promoter regions can be selected from any desired gene using CAT (chloramphenicol transferase) vectors or other vectors with selectable markers.

5 Two appropriate vectors are pKK232-8 and pCM7. Particular named bacterial promoters include lacI, lacZ, T3, T7, gpt, lambda PR, and trc. Eukaryotic promoters include CMV immediate early, HSV thymidine kinase, early and late SV40, LTRs from retrovirus, and mouse metallothionein- I. Selection of the appropriate vector and promoter is well within the level of ordinary skill in the art.

10 The present invention further provides host cells containing any one of the isolated fragments of the *Streptococcus pneumoniae* genomic fragments and contigs of the present invention, wherein the fragment has been introduced into the host cell using known methods. The host cell can be a higher eukaryotic host cell, such as a mammalian cell, a lower eukaryotic host cell, such as a yeast cell, or 15 a prokaryotic cell, such as a bacterial cell.

15 A polynucleotide of the present invention, such as a recombinant construct comprising an ORF of the present invention, may be introduced into the host by a variety of well established techniques that are standard in the art, such as calcium phosphate transfection, DEAE, dextran mediated transfection and electroporation, 20 which are described in, for instance, Davis, L. *et al.*, BASIC METHODS IN MOLECULAR BIOLOGY (1986).

25 A host cell containing one of the fragments of the *Streptococcus pneumoniae* genomic fragments and contigs of the present invention, can be used in conventional manners to produce the gene product encoded by the isolated fragment (in the case of an ORF) or can be used to produce a heterologous protein under the control of the EMF. The present invention further provides isolated polypeptides encoded by the nucleic acid fragments of the present invention or by degenerate variants of the nucleic acid fragments of the present invention. By "degenerate variant" is intended nucleotide fragments which differ 30 from a nucleic acid fragment of the present invention (e.g., an ORF) by nucleotide sequence but, due to the degeneracy of the Genetic Code, encode an identical polypeptide sequence.

Preferred nucleic acid fragments of the present invention are the ORFs and subfragments thereof depicted in Tables 2 and 3 which encode proteins.

A variety of methodologies known in the art can be utilized to obtain any one of the isolated polypeptides or proteins of the present invention. At the simplest level, the amino acid sequence can be synthesized using commercially available peptide synthesizers. This is particularly useful in producing small 5 peptides and fragments of larger polypeptides. Such short fragments as may be obtained most readily by synthesis are useful, for example, in generating antibodies against the native polypeptide, as discussed further below.

In an alternative method, the polypeptide or protein is purified from bacterial cells which naturally produce the polypeptide or protein. One skilled in 10 the art can readily employ well-known methods for isolating polypeptides and proteins to isolate and purify polypeptides or proteins of the present invention produced naturally by a bacterial strain, or by other methods. Methods for isolation and purification that can be employed in this regard include, but are not limited to, immunochromatography, HPLC, size-exclusion chromatography, ion-exchange chromatography, and immuno-affinity chromatography.

The polypeptides and proteins of the present invention also can be purified from cells which have been altered to express the desired polypeptide or protein. As used herein, a cell is said to be altered to express a desired polypeptide or protein when the cell, through genetic manipulation, is made to produce a 20 polypeptide or protein which it normally does not produce or which the cell normally produces at a lower level. Those skilled in the art can readily adapt procedures for introducing and expressing either recombinant or synthetic sequences into eukaryotic or prokaryotic cells in order to generate a cell which produces one of the polypeptides or proteins of the present invention.

25 Any host/vector system can be used to express one or more of the ORFs of the present invention. These include, but are not limited to, eukaryotic hosts such as HeLa cells, CV-1 cell, COS cells, and Sf9 cells, as well as prokaryotic host such as *E. coli* and *B. subtilis*. The most preferred cells are those which do not normally express the particular polypeptide or protein or which expresses the 30 polypeptide or protein at low natural level.

"Recombinant," as used herein, means that a polypeptide or protein is derived from recombinant (e.g., microbial or mammalian) expression systems. "Microbial" refers to recombinant polypeptides or proteins made in bacterial or fungal (e.g., yeast) expression systems. As a product, "recombinant microbial" defines a polypeptide or protein essentially free of native endogenous substances and unaccompanied by associated native glycosylation. Polypeptides or proteins expressed in most bacterial cultures, e.g., *E. coli*, will be free of glycosylation modifications; polypeptides or proteins expressed in yeast will have a glycosylation pattern different from that expressed in mammalian cells.

10 "Nucleotide sequence" refers to a heteropolymer of deoxyribonucleotides. Generally, DNA segments encoding the polypeptides and proteins provided by this invention are assembled from fragments of the *Streptococcus pneumoniae* genome and short oligonucleotide linkers, or from a series of oligonucleotides, to provide a synthetic gene which is capable of being expressed in a recombinant transcriptional 15 unit comprising regulatory elements derived from a microbial or viral operon.

20 "Recombinant expression vehicle or vector" refers to a plasmid or phage or virus or vector, for expressing a polypeptide from a DNA (RNA) sequence. The expression vehicle can comprise a transcriptional unit comprising an assembly of (1) a genetic regulatory elements necessary for gene expression in the host, 25 including elements required to initiate and maintain transcription at a level sufficient for suitable expression of the desired polypeptide, including, for example, promoters and, where necessary, an enhancer and a polyadenylation signal; (2) a structural or coding sequence which is transcribed into mRNA and translated into protein, and (3) appropriate signals to initiate translation at the beginning of the 30 desired coding region and terminate translation at its end. Structural units intended for use in yeast or eukaryotic expression systems preferably include a leader sequence enabling extracellular secretion of translated protein by a host cell. Alternatively, where recombinant protein is expressed without a leader or transport sequence, it may include an N-terminal methionine residue. This residue may or may not be subsequently cleaved from the expressed recombinant protein to provide a final product.

35 "Recombinant expression system" means host cells which have stably integrated a recombinant transcriptional unit into chromosomal DNA or carry the recombinant transcriptional unit extra chromosomally. The cells can be prokaryotic or eukaryotic. Recombinant expression systems as defined herein will express

heterologous polypeptides or proteins upon induction of the regulatory elements linked to the DNA segment or synthetic gene to be expressed.

5 Mature proteins can be expressed in mammalian cells, yeast, bacteria, or other cells under the control of appropriate promoters. Cell-free translation systems can also be employed to produce such proteins using RNAs derived from the DNA constructs of the present invention. Appropriate cloning and expression vectors for use with prokaryotic and eukaryotic hosts are described in Sambrook *et al.*, *Molecular Cloning: A Laboratory Manual*, 2nd Edition, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York (1989), the disclosure of which
10 is hereby incorporated by reference in its entirety.

15 Generally, recombinant expression vectors will include origins of replication and selectable markers permitting transformation of the host cell, *e.g.*, the ampicillin resistance gene of *E. coli* and *S. cerevisiae* TRP1 gene, and a promoter derived from a highly expressed gene to direct transcription of a downstream structural sequence. Such promoters can be derived from operons encoding glycolytic enzymes such as 3- phosphoglycerate kinase (PGK), alpha-factor, acid phosphatase, or heat shock proteins, among others. The heterologous structural sequence is assembled in appropriate phase with translation initiation and termination sequences, and preferably, a leader sequence capable of directing
20 secretion of translated protein into the periplasmic space or extracellular medium. Optionally, the heterologous sequence can encode a fusion protein including an N-terminal identification peptide imparting desired characteristics, *e.g.*, stabilization or simplified purification of expressed recombinant product.

25 Useful expression vectors for bacterial use are constructed by inserting a structural DNA sequence encoding a desired protein together with suitable translation initiation and termination signals in operable reading phase with a functional promoter. The vector will comprise one or more phenotypic selectable markers and an origin of replication to ensure maintenance of the vector and, when desirable, provide amplification within the host.

30 Suitable prokaryotic hosts for transformation include strains of *E. coli*, *B. subtilis*, *Salmonella typhimurium* and various species within the genera *Pseudomonas* and *Streptomyces*. Others may, also be employed as a matter of choice.

35 As a representative but non-limiting example, useful expression vectors for bacterial use can comprise a selectable marker and bacterial origin of replication

derived from commercially available plasmids comprising genetic elements of the well known cloning vector pBR322 (ATCC 37017). Such commercial vectors include, for example, pKK223-3 (available from Pharmacia Fine Chemicals, Uppsala, Sweden) and GEM 1 (available from Promega Biotec, Madison, WI, USA). These pBR322 "backbone" sections are combined with an appropriate promoter and the structural sequence to be expressed.

Following transformation of a suitable host strain and growth of the host strain to an appropriate cell density, the selected promoter, where it is inducible, is derepressed or induced by appropriate means (e.g., temperature shift or chemical induction) and cells are cultured for an additional period to provide for expression of the induced gene product. Thereafter cells are typically harvested, generally by centrifugation, disrupted to release expressed protein, generally by physical or chemical means, and the resulting crude extract is retained for further purification.

Various mammalian cell culture systems can also be employed to express recombinant protein. Examples of mammalian expression systems include the COS-7 lines of monkey kidney fibroblasts, described in Gluzman, *Cell* 23:175 (1981), and other cell lines capable of expressing a compatible vector, for example, the C127, 3T3, CHO, HeLa and BHK cell lines.

Mammalian expression vectors will comprise an origin of replication, a suitable promoter and enhancer, and also any necessary ribosome binding sites, polyadenylation site, splice donor and acceptor sites, transcriptional termination sequences, and 5' flanking nontranscribed sequences. DNA sequences derived from the SV40 viral genome, for example, SV40 origin, early promoter, enhancer, splice, and polyadenylation sites may be used to provide the required nontranscribed genetic elements.

Recombinant polypeptides and proteins produced in bacterial culture is usually isolated by initial extraction from cell pellets, followed by one or more salting-out, aqueous ion exchange or size exclusion chromatography steps. Microbial cells employed in expression of proteins can be disrupted by any convenient method, including freeze-thaw cycling, sonication, mechanical disruption, or use of cell lysing agents. Protein refolding steps can be used, as necessary, in completing configuration of the mature protein. Finally, high performance liquid chromatography (HPLC) can be employed for final purification steps.

The present invention further includes isolated polypeptides, proteins and nucleic acid molecules which are substantially equivalent to those herein described. As used herein, substantially equivalent can refer both to nucleic acid and amino acid sequences, for example a mutant sequence, that varies from a reference sequence by one or more substitutions, deletions, or additions, the net effect of which does not result in an adverse functional dissimilarity between reference and subject sequences. For purposes of the present invention, sequences having equivalent biological activity, and equivalent expression characteristics are considered substantially equivalent. For purposes of determining equivalence, 5 truncation of the mature sequence should be disregarded.

10

The invention further provides methods of obtaining homologs from other strains of *Streptococcus pneumoniae*, of the fragments of the *Streptococcus pneumoniae* genome of the present invention and homologs of the proteins encoded by the ORFs of the present invention. As used herein, a sequence or protein of 15 *Streptococcus pneumoniae* is defined as a homolog of a fragment of the *Streptococcus pneumoniae* fragments or contigs or a protein encoded by one of the ORFs of the present invention, if it shares significant homology to one of the fragments of the *Streptococcus pneumoniae* genome of the present invention or a protein encoded by one of the ORFs of the present invention. Specifically, by 20 using the sequence disclosed herein as a probe or as primers, and techniques such as PCR cloning and colony/plaque hybridization, one skilled in the art can obtain homologs.

As used herein, two nucleic acid molecules or proteins are said to "share significant homology" if the two contain regions which possess greater than 85% 25 sequence (amino acid or nucleic acid) homology. Preferred homologs in this regard are those with more than 90% homology. Especially preferred are those with 93% or more homology. Among especially preferred homologs those with 95% or more homology are particularly preferred. Very particularly preferred among these are those with 97% and even more particularly preferred among those 30 are homologs with 99% or more homology. The most preferred homologs among these are those with 99.9% homology or more. It will be understood that, among measures of homology, identity is particularly preferred in this regard.

Region specific primers or probes derived from the nucleotide sequence provided in SEQ ID NOS:1-391 or from a nucleotide sequence at least 95%, 35 particularly at least 99%, especially at least 99.5% identical to a sequence of SEQ

SEQ ID NOS:1-391 can be used to prime DNA synthesis and PCR amplification, as well as to identify colonies containing cloned DNA encoding a homolog. Methods suitable to this aspect of the present invention are well known and have been described in great detail in many publications such as, for example, Innis *et al.*, 5 *PCR Protocols*, Academic Press, San Diego, CA (1990)).

When using primers derived from SEQ ID NOS:1-391 or from a nucleotide sequence having an aforementioned identity to a sequence of SEQ ID NOS:1-391, one skilled in the art will recognize that by employing high stringency conditions (e.g., annealing at 50-60°C in 6X SSPC and 50% formamide, and washing at 50-10 65°C in 0.5X SSPC) only sequences which are greater than 75% homologous to the primer will be amplified. By employing lower stringency conditions (e.g., hybridizing at 35-37°C in 5X SSPC and 40-45% formamide, and washing at 42°C in 0.5X SSPC), sequences which are greater than 40-50% homologous to the primer will also be amplified.

15 When using DNA probes derived from SEQ ID NOS:1-391, or from a nucleotide sequence having an aforementioned identity to a sequence of SEQ ID NOS:1-391, for colony/plaque hybridization, one skilled in the art will recognize that by employing high stringency conditions (e.g., hybridizing at 50- 65°C in 5X SSPC and 50% formamide, and washing at 50- 65°C in 0.5X SSPC), sequences having regions which are greater than 90% homologous to the probe can be obtained, and that by employing lower stringency conditions (e.g., hybridizing at 20 35-37°C in 5X SSPC and 40-45% formamide, and washing at 42°C in 0.5X SSPC), sequences having regions which are greater than 35-45% homologous to the probe will be obtained.

25 Any organism can be used as the source for homologs of the present invention so long as the organism naturally expresses such a protein or contains genes encoding the same. The most preferred organism for isolating homologs are bacteria which are closely related to *Streptococcus pneumoniae*.

30 ILLUSTRATIVE USES OF COMPOSITIONS OF THE INVENTION

35 Each ORF provided in Tables 1 and 2 is identified with a function by homology to a known gene or polypeptide. As a result, one skilled in the art can use the polypeptides of the present invention for commercial, therapeutic and industrial purposes consistent with the type of putative identification of the

polypeptide. Such identifications permit one skilled in the art to use the *Streptococcus pneumoniae* ORFs in a manner similar to the known type of sequences for which the identification is made; for example, to ferment a particular sugar source or to produce a particular metabolite. A variety of reviews illustrative of this aspect of the invention are available, including the following reviews on the industrial use of enzymes, for example, BIOCHEMICAL ENGINEERING AND BIOTECHNOLOGY HANDBOOK, 2nd Ed., MacMillan Publications, Ltd. NY (1991) and BIOCATALYSTS IN ORGANIC SYNTHESSES, Tramper *et al.*, Eds., Elsevier Science Publishers, Amsterdam, The Netherlands (1985). A variety of exemplary uses that illustrate this and similar aspects of the present invention are discussed below.

1. Biosynthetic Enzymes

Open reading frames encoding proteins involved in mediating the catalytic reactions involved in intermediary and macromolecular metabolism, the biosynthesis of small molecules, cellular processes and other functions includes enzymes involved in the degradation of the intermediary products of metabolism, enzymes involved in central intermediary metabolism, enzymes involved in respiration, both aerobic and anaerobic, enzymes involved in fermentation, enzymes involved in ATP proton motor force conversion, enzymes involved in broad regulatory function, enzymes involved in amino acid synthesis, enzymes involved in nucleotide synthesis, enzymes involved in cofactor and vitamin synthesis, can be used for industrial biosynthesis.

The various metabolic pathways present in *Streptococcus pneumoniae* can be identified based on absolute nutritional requirements as well as by examining the various enzymes identified in Table 1-3 and SEQ ID NOS:1-391.

Of particular interest are polypeptides involved in the degradation of intermediary metabolites as well as non-macromolecular metabolism. Such enzymes include amylases, glucose oxidases, and catalase.

30 Proteolytic enzymes are another class of commercially important enzymes. Proteolytic enzymes find use in a number of industrial processes including the processing of flax and other vegetable fibers, in the extraction, clarification and depectinization of fruit juices, in the extraction of vegetables' oil and in the maceration of fruits and vegetables to give unicellular fruits. A detailed review of 35 the proteolytic enzymes used in the food industry is provided in Rombouts *et al.*,

Symbiosis 21:79 (1986) and Voragen *et al.* in *Biocatalysts In Agricultural Biotechnology*, Whitaker *et al.*, Eds., *American Chemical Society Symposium Series* 389:93 (1989).

The metabolism of sugars is an important aspect of the primary metabolism of *Streptococcus pneumoniae*. Enzymes involved in the degradation of sugars, such as, particularly, glucose, galactose, fructose and xylose, can be used in industrial fermentation. Some of the important sugar transforming enzymes, from a commercial viewpoint, include sugar isomerases such as glucose isomerase. Other metabolic enzymes have found commercial use such as glucose oxidases which produces ketogulonic acid (KGA). KGA is an intermediate in the commercial production of ascorbic acid using the Reichstein's procedure, as described in Krueger *et al.*, *Biotechnology* 6(A), Rhine *et al.*, Eds., Verlag Press, Weinheim, Germany (1984).

Glucose oxidase (GOD) is commercially available and has been used in purified form as well as in an immobilized form for the deoxygenation of beer. See, for instance, Hartmeir *et al.*, *Biotechnology Letters* 1:21 (1979). The most important application of GOD is the industrial scale fermentation of gluconic acid. Market for gluconic acids which are used in the detergent, textile, leather, photographic, pharmaceutical, food, feed and concrete industry, as described, for example, in Bigelis *et al.*, beginning on page 357 in *GENE MANIPULATIONS AND FUNGI*; Benett *et al.*, Eds., Academic Press, New York (1985). In addition to industrial applications, GOD has found applications in medicine for quantitative determination of glucose in body fluids recently in biotechnology for analyzing syrups from starch and cellulose hydrosylates. This application is described in Owusu *et al.*, *Biochem. et Biophysica. Acta.* 872:83 (1986), for instance.

The main sweetener used in the world today is sugar which comes from sugar beets and sugar cane. In the field of industrial enzymes, the glucose isomerase process shows the largest expansion in the market today. Initially, soluble enzymes were used and later immobilized enzymes were developed (Krueger *et al.*, *Biotechnology, The Textbook of Industrial Microbiology*, Sinauer Associated Incorporated, Sunderland, Massachusetts (1990)). Today, the use of glucose- produced high fructose syrups is by far the largest industrial business using immobilized enzymes. A review of the industrial use of these enzymes is provided by Jorgensen, *Starch* 40:307 (1988).

5 Proteinases, such as alkaline serine proteinases, are used as detergent additives and thus represent one of the largest volumes of microbial enzymes used in the industrial sector. Because of their industrial importance, there is a large body of published and unpublished information regarding the use of these enzymes in industrial processes. (See Faultman *et al.*, Acid Proteases Structure Function and Biology, Tang, J., ed., Plenum Press, New York (1977) and Godfrey *et al.*, Industrial Enzymes, MacMillan Publishers, Surrey, UK (1983) and Hepner *et al.*, Report Industrial Enzymes by 1990, Hel Hepner & Associates, London (1986)).

10 Another class of commercially usable proteins of the present invention are the microbial lipases, described by, for instance, Macrae *et al.*, *Philosophical Transactions of the Chiral Society of London* 310:227 (1985) and Poserke, *Journal of the American Oil Chemist Society* 61:1758 (1984). A major use of lipases is in the fat and oil industry for the production of neutral glycerides using lipase catalyzed inter-esterification of readily available triglycerides. Application of 15 lipases include the use as a detergent additive to facilitate the removal of fats from fabrics in the course of the washing procedures.

20 The use of enzymes, and in particular microbial enzymes, as catalyst for key steps in the synthesis of complex organic molecules is gaining popularity at a great rate. One area of great interest is the preparation of chiral intermediates. Preparation of chiral intermediates is of interest to a wide range of synthetic chemists particularly those scientists involved with the preparation of new pharmaceuticals, agrochemicals, fragrances and flavors. (See Davies *et al.*, *Recent 25 Advances in the Generation of Chiral Intermediates Using Enzymes*, CRC Press, Boca Raton, Florida (1990)). The following reactions catalyzed by enzymes are of interest to organic chemists: hydrolysis of carboxylic acid esters, phosphate esters, amides and nitriles, esterification reactions, trans-esterification reactions, synthesis of amides, reduction of alkanones and oxoalkanates, oxidation of alcohols to carbonyl compounds, oxidation of sulfides to sulfoxides, and carbon bond forming reactions such as the aldol reaction.

30 When considering the use of an enzyme encoded by one of the ORFs of the present invention for biotransformation and organic synthesis it is sometimes necessary to consider the respective advantages and disadvantages of using a microorganism as opposed to an isolated enzyme. Pros and cons of using a whole cell system on the one hand or an isolated partially purified enzyme on the other

hand, has been described in detail by Bud *et al.*, *Chemistry in Britain* (1987), p. 127.

5 Amino transferases, enzymes involved in the biosynthesis and metabolism of amino acids, are useful in the catalytic production of amino acids. The advantages of using microbial based enzyme systems is that the amino transferase enzymes catalyze the stereo-selective synthesis of only L-amino acids and generally possess uniformly high catalytic rates. A description of the use of amino transferases for amino acid production is provided by Roselle-David, *Methods of Enzymology* 136:479 (1987).

10 Another category of useful proteins encoded by the ORFs of the present invention include enzymes involved in nucleic acid synthesis, repair, and recombination.

2. Generation of Antibodies

15 As described here, the proteins of the present invention, as well as homologs thereof, can be used in a variety of procedures and methods known in the art which are currently applied to other proteins. The proteins of the present invention can further be used to generate an antibody which selectively binds the protein. Such antibodies can be either monoclonal or polyclonal antibodies, as well 20 fragments of these antibodies, and humanized forms.

25 The invention further provides antibodies which selectively bind to one of the proteins of the present invention and hybridomas which produce these antibodies. A hybridoma is an immortalized cell line which is capable of secreting a specific monoclonal antibody.

30 In general, techniques for preparing polyclonal and monoclonal antibodies as well as hybridomas capable of producing the desired antibody are well known in the art (Campbell, A. M., *Monoclonal Antibody Technology: Laboratory Techniques In Biochemistry And Molecular Biology*, Elsevier Science Publishers, Amsterdam, The Netherlands (1984); St. Groth *et al.*, *J. Immunol. Methods* 35: 1-21 (1980), Kohler and Milstein, *Nature* 256:495-497 (1975)), the trioma technique, the human B-cell hybridoma technique (Kozbor *et al.*, *Immunology Today* 4:72 (1983), pgs. 77-96 of Cole *et al.*, in *Monoclonal Antibodies And Cancer Therapy*, Alan R. Liss, Inc. (1985)). Any animal (mouse, rabbit, etc.) which is known to produce antibodies can be immunized with the pseudogene 35 polypeptide. Methods for immunization are well known in the art. Such methods

(1979); Engval, E. *et al.*, *Immunol.* 109:129 (1972); Goding, J. W., *J. Immunol. Meth.* 13:215 (1976)).

The labeled antibodies of the present invention can be used for *in vitro*, *in vivo*, and *in situ* assays to identify cells or tissues in which a fragment of the 5 *Streptococcus pneumoniae* genome is expressed.

The present invention further provides the above-described antibodies immobilized on a solid support. Examples of such solid supports include plastics such as polycarbonate, complex carbohydrates such as agarose and sepharose, acrylic resins and such as polyacrylamide and latex beads. Techniques for 10 coupling antibodies to such solid supports are well known in the art (Weir, D. M. *et al.*, "Handbook of Experimental Immunology" 4th Ed., Blackwell Scientific Publications, Oxford, England, Chapter 10 (1986); Jacoby, W. D. *et al.*, *Meth. Enzym.* 34 Academic Press, N. Y. (1974)). The immobilized antibodies of the 15 present invention can be used for *in vitro*, *in vivo*, and *in situ* assays as well as for immunoaffinity purification of the proteins of the present invention.

3. Diagnostic Assays and Kits

The present invention further provides methods to identify the expression of one of the ORFs of the present invention, or homolog thereof, in a test sample, 20 using one of the DFs or antibodies of the present invention.

In detail, such methods comprise incubating a test sample with one or more of the antibodies or one or more of the DFs of the present invention and assaying for binding of the DFs or antibodies to components within the test sample.

Conditions for incubating a DF or antibody with a test sample vary. 25 Incubation conditions depend on the format employed in the assay, the detection methods employed, and the type and nature of the DF or antibody used in the assay. One skilled in the art will recognize that any one of the commonly available hybridization, amplification or immunological assay formats can readily be adapted to employ the DFs or antibodies of the present invention. Examples of such assays 30 can be found in Chard, T., *An Introduction to Radioimmunoassay and Related Techniques*, Elsevier Science Publishers, Amsterdam, The Netherlands (1986); Bullock, G. R. *et al.*, *Techniques in Immunocytochemistry*, Academic Press, Orlando, FL Vol. 1 (1982), Vol. 2 (1983), Vol. 3 (1985); Tijssen, P., *Practice and Theory of Enzyme Immunoassays: Laboratory Techniques in Biochemistry and*

Molecular Biology, Elsevier Science Publishers, Amsterdam, The Netherlands (1985).

The test samples of the present invention include cells, protein or membrane extracts of cells, or biological fluids such as sputum, blood, serum, plasma, or urine. The test sample used in the above-described method will vary based on the assay format, nature of the detection method and the tissues, cells or extracts used as the sample to be assayed. Methods for preparing protein extracts or membrane extracts of cells are well known in the art and can be readily be adapted in order to obtain a sample which is compatible with the system utilized.

In another embodiment of the present invention, kits are provided which contain the necessary reagents to carry out the assays of the present invention.

Specifically, the invention provides a compartmentalized kit to receive, in close confinement, one or more containers which comprises: (a) a first container comprising one of the DFs or antibodies of the present invention; and (b) one or more other containers comprising one or more of the following: wash reagents, reagents capable of detecting presence of a bound DF or antibody.

In detail, a compartmentalized kit includes any kit in which reagents are contained in separate containers. Such containers include small glass containers, plastic containers or strips of plastic or paper. Such containers allows one to efficiently transfer reagents from one compartment to another compartment such that the samples and reagents are not cross-contaminated, and the agents or solutions of each container can be added in a quantitative fashion from one compartment to another. Such containers will include a container which will accept the test sample, a container which contains the antibodies used in the assay, containers which contain wash reagents (such as phosphate buffered saline, Tris-buffers, etc.), and containers which contain the reagents used to detect the bound antibody or DF.

Types of detection reagents include labelled nucleic acid probes, labelled secondary antibodies, or in the alternative, if the primary antibody is labelled, the enzymatic, or antibody binding reagents which are capable of reacting with the labelled antibody. One skilled in the art will readily recognize that the disclosed DFs and antibodies of the present invention can be readily incorporated into one of the established kit formats which are well known in the art.

4. Screening Assay for Binding Agents

Using the isolated proteins of the present invention, the present invention further provides methods of obtaining and identifying agents which bind to a protein encoded by one of the ORFs of the present invention or to one of the fragments and the *Streptococcus pneumoniae* fragment and contigs herein described.

5 In general, such methods comprise steps of:

(a) contacting an agent with an isolated protein encoded by one of the ORFs of the present invention, or an isolated fragment of the *Streptococcus pneumoniae* genome; and

10 (b) determining whether the agent binds to said protein or said fragment.

The agents screened in the above assay can be, but are not limited to, peptides, carbohydrates, vitamin derivatives, or other pharmaceutical agents. The agents can be selected and screened at random or rationally selected or designed using protein modeling techniques.

15 For random screening, agents such as peptides, carbohydrates, pharmaceutical agents and the like are selected at random and are assayed for their ability to bind to the protein encoded by the ORF of the present invention.

20 Alternatively, agents may be rationally selected or designed. As used herein, an agent is said to be "rationally selected or designed" when the agent is chosen based on the configuration of the particular protein. For example, one skilled in the art can readily adapt currently available procedures to generate peptides, pharmaceutical agents and the like capable of binding to a specific peptide sequence in order to generate rationally designed anti peptide peptides, for example see Hurby *et al.*, "Application of Synthetic Peptides: Antisense Peptides," in 25 *Synthetic Peptides, A User's Guide*, W. H. Freeman, NY (1992), pp. 289-307, and Kaspaczak *et al.*, *Biochemistry* 28:9230-8 (1989), or pharmaceutical agents, or the like.

30 In addition to the foregoing, one class of agents of the present invention, as broadly described, can be used to control gene expression through binding to one of the ORFs or EMFs of the present invention. As described above, such agents can be randomly screened or rationally designed/selected. Targeting the ORF or EMF allows a skilled artisan to design sequence specific or element specific agents, modulating the expression of either a single ORF or multiple ORFs which rely on the same EMF for expression control.

One class of DNA binding agents are agents which contain base residues which hybridize or form a triple helix by binding to DNA or RNA. Such agents can be based on the classic phosphodiester, ribonucleic acid backbone, or can be a variety of sulphydryl or polymeric derivatives which have base attachment capacity.

5 Agents suitable for use in these methods usually contain 20 to 40 bases and are designed to be complementary to a region of the gene involved in transcription (triple helix - see Lee *et al.*, *Nucl. Acids Res.* 6:3073 (1979); Cooney *et al.*, *Science* 241:456 (1988); and Dervan *et al.*, *Science* 251:1360 (1991)) or to the mRNA itself (antisense - Okano, *J. Neurochem.* 56:560 (1991);

10 10 *Oligodeoxynucleotides as Antisense Inhibitors of Gene Expression*, CRC Press, Boca Raton, FL (1988)). Triple helix- formation optimally results in a shut-off of RNA transcription from DNA, while antisense RNA hybridization blocks translation of an mRNA molecule into polypeptide. Both techniques have been demonstrated to be effective in model systems. Information contained in the

15 15 sequences of the present invention can be used to design antisense and triple helix-forming oligonucleotides, and other DNA binding agents.

5. Pharmaceutical Compositions and Vaccines

The present invention further provides pharmaceutical agents which can be used to modulate the growth or pathogenicity of *Streptococcus pneumoniae*, or another related organism, *in vivo* or *in vitro*. As used herein, a "pharmaceutical agent" is defined as a composition of matter which can be formulated using known techniques to provide a pharmaceutical compositions. As used herein, the "pharmaceutical agents of the present invention" refers the pharmaceutical agents which are derived from the proteins encoded by the ORFs of the present invention or are agents which are identified using the herein described assays.

As used herein, a pharmaceutical agent is said to "modulate the growth or pathogenicity of *Streptococcus pneumoniae* or a related organism, *in vivo* or *in vitro*," when the agent reduces the rate of growth, rate of division, or viability of the organism in question. The pharmaceutical agents of the present invention can modulate the growth or pathogenicity of an organism in many fashions, although an understanding of the underlying mechanism of action is not needed to practice the use of the pharmaceutical agents of the present invention. Some agents will modulate the growth by binding to an important protein thus blocking the biological activity of the protein, while other agents may bind to a component of the outer

surface of the organism blocking attachment or rendering the organism more prone to act the bodies nature immune system. Alternatively, the agent may comprise a protein encoded by one of the ORFs of the present invention and serve as a vaccine. The development and use of a vaccine based on outer membrane components are well known in the art.

5 As used herein, a "related organism" is a broad term which refers to any organism whose growth can be modulated by one of the pharmaceutical agents of the present invention. In general, such an organism will contain a homolog of the protein which is the target of the pharmaceutical agent or the protein used as a vaccine. As such, related organisms do not need to be bacterial but may be fungal 10 or viral pathogens.

10 The pharmaceutical agents and compositions of the present invention may be administered in a convenient manner, such as by the oral, topical, intravenous, intraperitoneal, intramuscular, subcutaneous, intranasal or intradermal routes. The 15 pharmaceutical compositions are administered in an amount which is effective for treating and/or prophylaxis of the specific indication. In general, they are administered in an amount of at least about 1 mg/kg body weight and in most cases they will be administered in an amount not in excess of about 1 g/kg body weight per day. In most cases, the dosage is from about 0.1 mg/kg to about 10 g/kg body 20 weight daily, taking into account the routes of administration, symptoms, etc.

20 The agents of the present invention can be used in native form or can be modified to form a chemical derivative. As used herein, a molecule is said to be a "chemical derivative" of another molecule when it contains additional chemical 25 moieties not normally a part of the molecule. Such moieties may improve the molecule's solubility, absorption, biological half life, etc. The moieties may alternatively decrease the toxicity of the molecule, eliminate or attenuate any undesirable side effect of the molecule, etc. Moieties capable of mediating such effects are disclosed in, among other sources, REMINGTON'S PHARMACEUTICAL SCIENCES (1980) cited elsewhere herein.

30 For example, such moieties may change an immunological character of the functional derivative, such as affinity for a given antibody. Such changes in immunomodulation activity are measured by the appropriate assay, such as a competitive type immunoassay. Modifications of such protein properties as redox or thermal stability, biological half-life, hydrophobicity, susceptibility to proteolytic 35 degradation or the tendency to aggregate with carriers or into multimers also may

be effected in this way and can be assayed by methods well known to the skilled artisan.

The therapeutic effects of the agents of the present invention may be obtained by providing the agent to a patient by any suitable means (*e.g.*, inhalation, 5 intravenously, intramuscularly, subcutaneously, enterally, or parenterally). It is preferred to administer the agent of the present invention so as to achieve an effective concentration within the blood or tissue in which the growth of the organism is to be controlled. To achieve an effective blood concentration, the preferred method is to administer the agent by injection. The administration may be 10 by continuous infusion, or by single or multiple injections.

In providing a patient with one of the agents of the present invention, the dosage of the administered agent will vary depending upon such factors as the patient's age, weight, height, sex, general medical condition, previous medical history, *etc.* In general, it is desirable to provide the recipient with a dosage of 15 agent which is in the range of from about 1 pg/kg to 10 mg/kg (body weight of patient), although a lower or higher dosage may be administered. The therapeutically effective dose can be lowered by using combinations of the agents of the present invention or another agent.

As used herein, two or more compounds or agents are said to be 20 administered "in combination" with each other when either (1) the physiological effects of each compound, or (2) the serum concentrations of each compound can be measured at the same time. The composition of the present invention can be administered concurrently with, prior to, or following the administration of the other agent.

25 The agents of the present invention are intended to be provided to recipient subjects in an amount sufficient to decrease the rate of growth (as defined above) of the target organism.

The administration of the agent(s) of the invention may be for either a 30 "prophylactic" or "therapeutic" purpose. When provided prophylactically, the agent(s) are provided in advance of any symptoms indicative of the organisms growth. The prophylactic administration of the agent(s) serves to prevent, attenuate, or decrease the rate of onset of any subsequent infection. When provided therapeutically, the agent(s) are provided at (or shortly after) the onset of an indication of infection. The therapeutic administration of the compound(s)

serves to attenuate the pathological symptoms of the infection and to increase the rate of recovery.

The agents of the present invention are administered to a subject, such as a mammal, or a patient, in a pharmaceutically acceptable form and in a therapeutically effective concentration. A composition is said to be "pharmacologically acceptable" if its administration can be tolerated by a recipient patient. Such an agent is said to be administered in a "therapeutically effective amount" if the amount administered is physiologically significant. An agent is physiologically significant if its presence results in a detectable change in the physiology of a recipient patient.

The agents of the present invention can be formulated according to known methods to prepare pharmaceutically useful compositions, whereby these materials, or their functional derivatives, are combined in a mixture with a pharmaceutically acceptable carrier vehicle. Suitable vehicles and their formulation, inclusive of other human proteins, e.g., human serum albumin, are described, for example, in REMINGTON'S PHARMACEUTICAL SCIENCES, 16th Ed., Osol, A., Ed., Mack Publishing, Easton PA (1980). In order to form a pharmaceutically acceptable composition suitable for effective administration, such compositions will contain an effective amount of one or more of the agents of the present invention, together with a suitable amount of carrier vehicle.

Additional pharmaceutical methods may be employed to control the duration of action. Control release preparations may be achieved through the use of polymers to complex or absorb one or more of the agents of the present invention. The controlled delivery may be effectuated by a variety of well known techniques, including formulation with macromolecules such as, for example, polyesters, polyamino acids, polyvinyl, pyrrolidone, ethylenevinylacetate, methylcellulose, carboxymethylcellulose, or protamine, sulfate, adjusting the concentration of the macromolecules and the agent in the formulation, and by appropriate use of methods of incorporation, which can be manipulated to effectuate a desired time course of release. Another possible method to control the duration of action by controlled release preparations is to incorporate agents of the present invention into particles of a polymeric material such as polyesters, polyamino acids, hydrogels, poly(lactic acid) or ethylene vinylacetate copolymers. Alternatively, instead of incorporating these agents into polymeric particles, it is possible to entrap these materials in microcapsules prepared, for example, by coacervation techniques or by interfacial polymerization with, for example, hydroxymethylcellulose or gelatine-

microcapsules and poly(methylmethacrylate) microcapsules, respectively, or in colloidal drug delivery systems, for example, liposomes, albumin microspheres, microemulsions, nanoparticles, and nanocapsules or in macroemulsions. Such techniques are disclosed in REMINGTON'S PHARMACEUTICAL SCIENCES

5 (1980).

The invention further provides a pharmaceutical pack or kit comprising one or more containers filled with one or more of the ingredients of the pharmaceutical compositions of the invention. Associated with such container(s) can be a notice in the form prescribed by a governmental agency regulating the manufacture, use or 10 sale of pharmaceuticals or biological products, which notice reflects approval by the agency of manufacture, use or sale for human administration.

In addition, the agents of the present invention may be employed in conjunction with other therapeutic compounds.

15 6. Shot-Gun Approach to Megabase DNA Sequencing

The present invention further demonstrates that a large sequence can be sequenced using a random shotgun approach. This procedure, described in detail in the examples that follow, has eliminated the up front cost of isolating and ordering overlapping or contiguous subclones prior to the start of the sequencing 20 protocols.

Certain aspects of the present invention are described in greater detail in the examples that follow. The examples are provided by way of illustration. Other aspects and embodiments of the present invention are contemplated by the inventors, as will be clear to those of skill in the art from reading the present 25 disclosure.

ILLUSTRATIVE EXAMPLES

LIBRARIES AND SEQUENCING

1. Shotgun Sequencing Probability Analysis

30 The overall strategy for a shotgun approach to whole genome sequencing follows from the Lander and Waterman (Landerman and Waterman, *Genomics* 2:231 (1988)) application of the equation for the Poisson distribution. According to this treatment, the probability, P_0 , that any given base in a sequence of size L , in 35 nucleotides, is not sequenced after a certain amount, n , in nucleotides, of random

sequence has been determined can be calculated by the equation $P = e^{-m}$, where m is L/n , the fold coverage. For instance, for a genome of 2.8 Mb, $m=1$ when 2.8 Mb of sequence has been randomly generated (1X coverage). At that point, $P = e^{-1} = 0.37$. The probability that any given base has not been sequenced is the same as the probability that any region of the whole sequence L has not been determined and, therefore, is equivalent to the fraction of the whole sequence that has yet to be determined. Thus, at one-fold coverage, approximately 37% of a polynucleotide of size L , in nucleotides has not been sequenced. When 14 Mb of sequence has been generated, coverage is 5X for a 2.8 Mb and the unsequenced fraction drops to .0067 or 0.67%. 5X coverage of a 2.8 Mb sequence can be attained by sequencing approximately 17,000 random clones from both insert ends with an average sequence read length of 410 bp.

Similarly, the total gap length, G , is determined by the equation $G = Le^{-m}$, and the average gap size, g , follows the equation, $g = L/n$. Thus, 5X coverage leaves about 240 gaps averaging about 82 bp in size in a sequence of a polynucleotide 2.8 Mb long.

The treatment above is essentially that of Lander and Waterman, *Genomics* 2: 231 (1988).

20 2. Random Library Construction

In order to approximate the random model described above during actual sequencing, a nearly ideal library of cloned genomic fragments is required. The following library construction procedure was developed to achieve this end.

25 *Streptococcus pneumoniae* DNA is prepared by phenol extraction. A mixture containing 200 μ g DNA in 1.0 ml of 300 mM sodium acetate, 10 mM Tris-HCl, 1 mM Na-EDTA, 50% glycerol is processed through a nebulizer (IPI Medical Products) with a stream of nitrogen adjusted to 35 Kpa for 2 minutes. The sonicated DNA is ethanol precipitated and redissolved in 500 μ l TE buffer.

To create blunt-ends, a 100 μ l aliquot of the resuspended DNA is digested 30 with 5 units of BAL31 nuclease (New England BioLabs) for 10 min at 30°C in 200 μ l BAL31 buffer. The digested DNA is phenol-extracted, ethanol-precipitated, redissolved in 100 μ l TE buffer, and then size-fractionated by electrophoresis through a 1.0% low melting temperature agarose gel. The section containing DNA fragments 1.6-2.0 kb in size is excised from the gel, and the LGT agarose is melted 35 and the resulting solution is extracted with phenol to separate the agarose from the

DNA. DNA is ethanol precipitated and redissolved in 20 μ l of TE buffer for ligation to vector.

A two-step ligation procedure is used to produce a plasmid library with 97% inserts, of which >99% were single inserts. The first ligation mixture (50 μ l) 5 contains 2 μ g of DNA fragments, 2 μ g pUC18 DNA (Pharmacia) cut with SmaI and dephosphorylated with bacterial alkaline phosphatase, and 10 units of T4 ligase (GIBCO/BRL) and is incubated at 14°C for 4 hr. The ligation mixture then is phenol extracted and ethanol precipitated, and the precipitated DNA is dissolved in 20 μ l TE buffer and electrophoresed on a 1.0% low melting agarose gel. Discrete 10 bands in a ladder are visualized by ethidium bromide-staining and UV illumination and identified by size as insert (I), vector (v), v+I, v+2i, v+3i, etc. The portion of the gel containing v+I DNA is excised and the v+I DNA is recovered and resuspended into 20 μ l TE. The v+I DNA then is blunt-ended by T4 polymerase treatment for 5 min. at 37°C in a reaction mixture (50 μ l) containing the v+I linears, 15 500 μ M each of the 4 dNTPs, and 9 units of T4 polymerase (New England BioLabs), under recommended buffer conditions. After phenol extraction and ethanol precipitation the repaired v+I linears are dissolved in 20 μ l TE. The final ligation to produce circles is carried out in a 50 μ l reaction containing 5 μ l of v+I 20 linears and 5 units of T4 ligase at 14°C overnight. After 10 min. at 70°C the following day, the reaction mixture is stored at -20°C.

This two-stage procedure results in a molecularly random collection of single-insert plasmid recombinants with minimal contamination from double-insert chimeras (<1%) or free vector (<3%).

Since deviation from randomness can arise from propagation the DNA in 25 the host, *E. coli* host cells deficient in all recombination and restriction functions (A. Greener, *Strategies* 3 (1):5 (1990)) are used to prevent rearrangements, deletions, and loss of clones by restriction. Furthermore, transformed cells are plated directly on antibiotic diffusion plates to avoid the usual broth recovery phase which allows multiplication and selection of the most rapidly growing cells.

30 Plating is carried out as follows. A 100 μ l aliquot of Epicurian Coli SURE II Supercompetent Cells (Stratagene 200152) is thawed on ice and transferred to a chilled Falcon 2059 tube on ice. A 1.7 μ l aliquot of 1.42 M beta-mercaptoethanol is added to the aliquot of cells to a final concentration of 25 mM. Cells are incubated on ice for 10 min. A 1 μ l aliquot of the final ligation is added to the cells 35 and incubated on ice for 30 min. The cells are heat pulsed for 30 sec. at 42°C and

placed back on ice for 2 min. The outgrowth period in liquid culture is eliminated from this protocol in order to minimize the preferential growth of any given transformed cell. Instead the transformation mixture is plated directly on a nutrient rich SOB plate containing a 5 ml bottom layer of SOB agar (5% SOB agar: 20 g 5 tryptone, 5 g yeast extract, 0.5 g NaCl, 1.5% Difco Agar per liter of media). The 5 ml bottom layer is supplemented with 0.4 ml of 50 mg/ml ampicillin per 100 ml SOB agar. The 15 ml top layer of SOB agar is supplemented with 1 ml X-Gal (2%), 1 ml MgCl (1 M), and 1 ml MgSO /100 ml SOB agar. The 15 ml top layer is poured just prior to plating. Our titer is approximately 100 colonies/10 μ l aliquot 10 of transformation.²

All colonies are picked for template preparation regardless of size. Thus, only clones lost due to "poison" DNA or deleterious gene products are deleted from the library, resulting in a slight increase in gap number over that expected.

15 3. Random DNA Sequencing

High quality double stranded DNA plasmid templates are prepared using a "boiling bead" method developed in collaboration with Advanced Genetic Technology Corp. (Gaithersburg, MD) (Adams *et al.*, *Science* 252:1651 (1991); Adams *et al.*, *Nature* 355:632 (1992)). Plasmid preparation is performed in a 96-well format for all stages of DNA preparation from bacterial growth through final DNA purification. Template concentration is determined using Hoechst Dye and a Millipore Cytofluor. DNA concentrations are not adjusted, but low-yielding templates are identified where possible and not sequenced.

Templates are also prepared from two *Streptococcus pneumoniae* lambda 25 genomic libraries. An amplified library is constructed in the vector Lambda GEM-12 (Promega) and an unamplified library is constructed in Lambda DASH II (Stratagene). In particular, for the unamplified lambda library, *Streptococcus pneumoniae* DNA (> 100 kb) is partially digested in a reaction mixture (200 μ l) containing 50 μ g DNA, 1X Sau3AI buffer, 20 units Sau3AI for 6 min. at 23°C. 30 The digested DNA was phenol-extracted and electrophoresed on a 0.5% low melting agarose gel at 2V/cm for 7 hours. Fragments from 15 to 25 kb are excised and recovered in a final volume of 6 μ l. One μ l of fragments is used with 1 μ l of DASHII vector (Stratagene) in the recommended ligation reaction. One μ l of the ligation mixture is used per packaging reaction following the recommended 35 protocol with the Gigapack II XL Packaging Extract (Stratagene, #227711). Phage

are plated directly without amplification from the packaging mixture (after dilution with 500 μ l of recommended SM buffer and chloroform treatment). Yield is about 2.5×10^3 pfu/ μ l. The amplified library is prepared essentially as above except the lambda GEM-12 vector is used. After packaging, about 3.5×10^4 pfu are plated on 5 the restrictive NM539 host. The lysate is harvested in 2 ml of SM buffer and stored frozen in 7% dimethylsulfoxide. The phage titer is approximately 1×10^9 pfu/ml.

10 Liquid lysates (100 μ l) are prepared from randomly selected plaques (from the unamplified library) and template is prepared by long-range PCR using T7 and T3 vector-specific primers.

15 Sequencing reactions are carried out on plasmid and/or PCR templates using the AB Catalyst LabStation with Applied Biosystems PRISM Ready Reaction Dye Primer Cycle Sequencing Kits for the M13 forward (M13-21) and the M13 reverse (M13RP1) primers (Adams *et al.*, *Nature* 368:474 (1994)). Dye terminator sequencing reactions are carried out on the lambda templates on a Perkin-Elmer 9600 Thermocycler using the Applied Biosystems Ready Reaction Dye Terminator Cycle Sequencing kits. T7 and SP6 primers are used to sequence the ends of the inserts from the Lambda GEM-12 library and T7 and T3 primers are used to sequence the ends of the inserts from the Lambda DASH II library.

20 Sequencing reactions are performed by eight individuals using an average of fourteen AB 373 DNA Sequencers per day. All sequencing reactions are analyzed using the Stretch modification of the AB 373, primarily using a 34 cm well-to-read distance. The overall sequencing success rate very approximately is about 85% for M13-21 and M13RP1 sequences and 65% for dye-terminator reactions. The 25 average usable read length is 485 bp for M13-21 sequences, 445bp for M13RP1 sequences, and 375 bp for dye-terminator reactions.

30 Richards *et al.*, Chapter 28 in AUTOMATED DNA SEQUENCING AND ANALYSIS, M. D. Adams, C. Fields, J. C. Venter, Eds., Academic Press, London, (1994) described the value of using sequence from both ends of sequencing templates to facilitate ordering of contigs in shotgun assembly projects of lambda and cosmid clones. We balance the desirability of both-end sequencing (including the reduced cost of lower total number of templates) against shorter 35 read-lengths for sequencing reactions performed with the M13RP1 (reverse) primer compared to the M13-21 (forward) primer. Approximately one-half of the templates are sequenced from both ends. Random reverse sequencing reactions are

done based on successful forward sequencing reactions. Some M13RP1 sequences are obtained in a semi-directed fashion: M13-21: sequences pointing outward at the ends of contigs are chosen for M13RP1 sequencing in an effort to specifically order contigs.

5

4. Protocol for Automated Cycle Sequencing

The sequencing is carried out using ABI Catalyst robots and AB 373 Automated DNA Sequencers. The Catalyst robot is a publicly available sophisticated pipetting and temperature control robot which has been developed 10 specifically for DNA sequencing reactions. The Catalyst combines pre-aliquoted templates and reaction mixes consisting of deoxy- and dideoxynucleotides, the thermostable Taq DNA polymerase, fluorescently-labelled sequencing primers, and reaction buffer. Reaction mixes and templates are combined in the wells of an aluminum 96-well thermocycling plate. Thirty consecutive cycles of linear 15 amplification (*i.e.*, one primer synthesis) steps are performed including denaturation, annealing of primer and template, and extension; *i.e.*, DNA synthesis. A heated lid with rubber gaskets on the thermocycling plate prevents evaporation without the need for an oil overlay.

Two sequencing protocols are used: one for dye-labelled primers and a 20 second for dye-labelled dideoxy chain terminators. The shotgun sequencing involves use of four dye-labelled sequencing primers, one for each of the four terminator nucleotide. Each dye-primer is labelled with a different fluorescent dye, permitting the four individual reactions to be combined into one lane of the 373 DNA Sequencer for electrophoresis, detection, and base-calling. ABI currently 25 supplies pre-mixed reaction mixes in bulk packages containing all the necessary non-template reagents for sequencing. Sequencing can be done with both plasmid and PCR- generated templates with both dye-primers and dye- terminators with approximately equal fidelity, although plasmid templates generally give longer usable sequences.

30

Thirty-two reactions are loaded per AB373 Sequencer each day, for a total of 960 samples. Electrophoresis is run overnight following the manufacturer's protocols, and the data is collected for twelve hours. Following electrophoresis and fluorescence detection, the ABI 373 performs automatic lane tracking and base-calling. The lane-tracking is confirmed visually. Each sequence electropherogram 35 (or fluorescence lane trace) is inspected visually and assessed for quality. Trailing

sequences of low quality are removed and the sequence itself is loaded via software to a Sybase database (archived daily to 8mm tape). Leading vector polylinker sequence is removed automatically by a software program. Average edited lengths of sequences from the standard ABI 373 are around 400 bp and depend mostly on 5 the quality of the template used for the sequencing reaction. ABI 373 Sequencers converted to Stretch Liners provide a longer electrophoresis path prior to fluorescence detection and increase the average number of usable bases to 500-600 bp.

10 **INFORMATICS**

1. Data Management

A number of information management systems for a large-scale sequencing lab have been developed. (For review see, for instance, Kerlavage *et al.*, *Proceedings of the Twenty-Sixth Annual Hawaii International Conference on System Sciences*, IEEE Computer Society Press, Washington D. C., 585 (1993)) 15 The system used to collect and assemble the sequence data was developed using the Sybase relational database management system and was designed to automate data flow wherever possible and to reduce user error. The database stores and correlates all information collected during the entire operation from template 20 preparation to final analysis of the genome. Because the raw output of the ABI 373 Sequencers was based on a Macintosh platform and the data management system chosen was based on a Unix platform, it was necessary to design and implement a variety of multi- user, client-server applications which allow the raw data as well as analysis results to flow seamlessly into the database with a minimum of user effort.

25

2. Assembly

An assembly engine (TIGR Assembler) developed for the rapid and accurate assembly of thousands of sequence fragments was employed to generate contigs. The TIGR assembler simultaneously clusters and assembles fragments of 30 the genome. In order to obtain the speed necessary to assemble more than 10^4 fragments, the algorithm builds a hash table of 12 bp oligonucleotide subsequences to generate a list of potential sequence fragment overlaps. The number of potential overlaps for each fragment determines which fragments are likely to fall into repetitive elements. Beginning with a single seed sequence fragment, TIGR 35 Assembler extends the current contig by attempting to add the best matching

fragment based on oligonucleotide content. The contig and candidate fragment are aligned using a modified version of the Smith-Waterman algorithm which provides for optimal gapped alignments (Waterman, M. S., *Methods in Enzymology* 164:765 (1988)). The contig is extended by the fragment only if strict criteria for 5 the quality of the match are met. The match criteria include the minimum length of overlap, the maximum length of an unmatched end, and the minimum percentage match. These criteria are automatically lowered by the algorithm in regions of minimal coverage and raised in regions with a possible repetitive element. The 10 number of potential overlaps for each fragment determines which fragments are likely to fall into repetitive elements. Fragments representing the boundaries of repetitive elements and potentially chimeric fragments are often rejected based on partial mismatches at the ends of alignments and excluded from the current contig. TIGR Assembler is designed to take advantage of clone size information coupled 15 with sequencing from both ends of each template. It enforces the constraint that sequence fragments from two ends of the same template point toward one another in the contig and are located within a certain range of base pairs (definable for each clone based on the known clone size range for a given library).

The process resulted in 391 contigs as represented by SEQ ID NOs:1-391.

20 3. Identifying Genes

The predicted coding regions of the *Streptococcus pneumoniae* genome were initially defined with the program GeneMark, which finds ORFs using a probabilistic classification technique. The predicted coding region sequences were used in searches against a database of all nucleotide sequences from GenBank 25 (October, 1997), using the BLASTN search method to identify overlaps of 50 or more nucleotides with at least a 95% identity. Those ORFs with nucleotide sequence matches are shown in Table 1. The ORFs without such matches were translated to protein sequences and compared to a non-redundant database of known proteins generated by combining the Swiss-prot, PIR and GenPept 30 databases. ORFs that matched a database protein with BLASTP probability less than or equal to 0.01 are shown in Table 2. The table also lists assigned functions based on the closest match in the databases. ORFs that did not match protein or nucleotide sequences in the databases at these levels are shown in Table 3.

ILLUSTRATIVE APPLICATIONS

1. Production of an Antibody to a *Streptococcus pneumoniae* Protein

Substantially pure protein or polypeptide is isolated from the transfected or 5 transformed cells using any one of the methods known in the art. The protein can also be produced in a recombinant prokaryotic expression system, such as *E. coli*, or can be chemically synthesized. Concentration of protein in the final preparation is adjusted, for example, by concentration on an Amicon filter device, to the level of a few micrograms/ml. Monoclonal or polyclonal antibody to the protein can 10 then be prepared as follows.

2. Monoclonal Antibody Production by Hybridoma Fusion

Monoclonal antibody to epitopes of any of the peptides identified and isolated as described can be prepared from murine hybridomas according to the 15 classical method of Kohler, G. and Milstein, C., *Nature* 256:495 (1975) or modifications of the methods thereof. Briefly, a mouse is repetitively inoculated with a few micrograms of the selected protein over a period of a few weeks. The mouse is then sacrificed, and the antibody producing cells of the spleen isolated. The spleen cells are fused by means of polyethylene glycol with mouse myeloma 20 cells, and the excess unfused cells destroyed by growth of the system on selective media comprising aminopterin (HAT media). The successfully fused cells are diluted and aliquots of the dilution placed in wells of a microtiter plate where growth of the culture is continued. Antibody-producing clones are identified by detection of antibody in the supernatant fluid of the wells by immunoassay 25 procedures, such as ELISA, as originally described by Engvall, E., *Meth. Enzymol.* 70:419 (1980), and modified methods thereof. Selected positive clones can be expanded and their monoclonal antibody product harvested for use. Detailed procedures for monoclonal antibody production are described in Davis, L. *et al.*, *Basic Methods in Molecular Biology*, Elsevier, New York. Section 21-2 (1989).

3. Polyclonal Antibody Production by Immunization

5 Polyclonal antiserum containing antibodies to heterogenous epitopes of a single protein can be prepared by immunizing suitable animals with the expressed protein described above, which can be unmodified or modified to enhance immunogenicity. Effective polyclonal antibody production is affected by many factors related both to the antigen and the host species. For example, small molecules tend to be less immunogenic than others and may require the use of carriers and adjuvant. Also, host animals vary in response to site of inoculations and dose, with both inadequate or excessive doses of antigen resulting in low titer 10 antisera. Small doses (ng level) of antigen administered at multiple intradermal sites appears to be most reliable. An effective immunization protocol for rabbits can be found in Vaitukaitis, J. *et al.*, *J. Clin. Endocrinol. Metab.* 33:988-991 (1971).

15 Booster injections can be given at regular intervals, and antiserum harvested when antibody titer thereof, as determined semi-quantitatively, for example, by double immunodiffusion in agar against known concentrations of the antigen, begins to fall. See, for example, Ouchterlony, O. *et al.*, Chap. 19 in: *Handbook of Experimental Immunology*, Wier, D., ed, Blackwell (1973). Plateau concentration of antibody is usually in the range of 0.1 to 0.2 mg/ml of serum (about 12M). 20 Affinity of the antisera for the antigen is determined by preparing competitive binding curves, as described, for example, by Fisher, D., Chap. 42 in: *Manual of Clinical Immunology*, second edition, Rose and Friedman, eds., Amer. Soc. For Microbiology, Washington, D. C. (1980)

25 Antibody preparations prepared according to either protocol are useful in quantitative immunoassays which determine concentrations of antigen-bearing substances in biological samples; they are also used semi- quantitatively or qualitatively to identify the presence of antigen in a biological sample. In addition, antibodies are useful in various animal models of pneumococcal disease as a means of evaluating the protein used to make the antibody as a potential vaccine target or 30 as a means of evaluating the antibody as a potential immunotherapeutic or immunoprophylactic reagent.

4. Preparation of PCR Primers and Amplification of DNA

Various fragments of the *Streptococcus pneumoniae* genome, such as those of Tables 1-3 and SEQ ID NOS:1-391 can be used, in accordance with the present invention, to prepare PCR primers for a variety of uses. The PCR primers are 5 preferably at least 15 bases, and more preferably at least 18 bases in length. When selecting a primer sequence, it is preferred that the primer pairs have approximately the same G/C ratio, so that melting temperatures are approximately the same. The PCR primers and amplified DNA of this Example find use in the Examples that follow.

10

5. Gene expression from DNA Sequences Corresponding to ORFs

A fragment of the *Streptococcus pneumoniae* genome provided in Tables 1-3 is introduced into an expression vector using conventional technology. 15 Techniques to transfer cloned sequences into expression vectors that direct protein translation in mammalian, yeast, insect or bacterial expression systems are well known in the art. Commercially available vectors and expression systems are available from a variety of suppliers including Stratagene (La Jolla, California), Promega (Madison, Wisconsin), and Invitrogen (San Diego, California). If 20 desired, to enhance expression and facilitate proper protein folding, the codon context and codon pairing of the sequence may be optimized for the particular expression organism, as explained by Hatfield *et al.*, U. S. Patent No. 5,082,767, incorporated herein by this reference.

The following is provided as one exemplary method to generate polypeptide(s) from cloned ORFs of the *Streptococcus pneumoniae* genome fragment. Bacterial ORFs generally lack a poly A addition signal. The addition signal sequence can be added to the construct by, for example, splicing out the poly 5 A addition sequence from pSG5 (Stratagene) using BglII and SalI restriction endonuclease enzymes and incorporating it into the mammalian expression vector pXT1 (Stratagene) for use in eukaryotic expression systems. pXT1 contains the LTRs and a portion of the gag gene of Moloney Murine Leukemia Virus. The positions of the LTRs in the construct allow efficient stable transfection. The 10 vector includes the Herpes Simplex thymidine kinase promoter and the selectable neomycin gene. The *Streptococcus pneumoniae* DNA is obtained by PCR from the bacterial vector using oligonucleotide primers complementary to the *Streptococcus pneumoniae* DNA and containing restriction endonuclease sequences for PstI incorporated into the 5' primer and BglII at the 5' end of the corresponding 15 *Streptococcus pneumoniae* DNA 3' primer, taking care to ensure that the *Streptococcus pneumoniae* DNA is positioned such that it is followed with the poly A addition sequence. The purified fragment obtained from the resulting PCR reaction is digested with PstI, blunt ended with an exonuclease, digested with BglII, purified and ligated to pXT1, now containing a poly A addition sequence 20 and digested BglII.

The ligated product is transfected into mouse NIH 3T3 cells using Lipofectin (Life Technologies, Inc., Grand Island, New York) under conditions outlined in the product specification. Positive transfectants are selected after growing the transfected cells in 600 ug/ml G418 (Sigma, St. Louis, Missouri). 25 The protein is preferably released into the supernatant. However if the protein has membrane binding domains, the protein may additionally be retained within the cell or expression may be restricted to the cell surface. Since it may be necessary to purify and locate the transfected product, synthetic 15-mer peptides synthesized from the predicted *Streptococcus pneumoniae* DNA sequence are injected into mice 30 to generate antibody to the polypeptide encoded by the *Streptococcus pneumoniae* DNA.

Alternatively and if antibody production is not possible, the *Streptococcus pneumoniae* DNA sequence is additionally incorporated into eukaryotic expression vectors and expressed as, for example, a globin fusion. Antibody to the globin moiety then is used to purify the chimeric protein. Corresponding protease 5 cleavage sites are engineered between the globin moiety and the polypeptide encoded by the *Streptococcus pneumoniae* DNA so that the latter may be freed from the formed by simple protease digestion. One useful expression vector for generating globin chimerics is pSG5 (Stratagene). This vector encodes a rabbit 10 globin. Intron II of the rabbit globin gene facilitates splicing of the expressed transcript, and the polyadenylation signal incorporated into the construct increases the level of expression. These techniques are well known to those skilled in the art of molecular biology. Standard methods are published in methods texts such as Davis *et al.*, cited elsewhere herein, and many of the methods are available from the 15 technical assistance representatives from Stratagene, Life Technologies, Inc., or Promega. Polypeptides of the invention also may be produced using *in vitro* translation systems such as *in vitro* ExpressTM Translation Kit (Stratagene).

While the present invention has been described in some detail for purposes of clarity and understanding, one skilled in the art will appreciate that various changes in form and detail can be made without departing from the true scope of the invention.

All patents, patent applications and publications referred to above are hereby incorporated by reference.

TABLE 1 *S. pneumoniae* - Coding regions containing known sequences

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	percent HSP nt ident	HSP nt length	ORF nt length
1	1	437	1003	gb U41755	Streptococcus pneumoniae peptide methionine sulfoxide reductase (msrA) and homoserine kinase homolog (thrB) genes, complete cds	92	200	567
2	5	6169	5720	gb U004047	Streptococcus pneumoniae S52 dextran glucosidase gene and insertion sequence IS1202 transposase gene, complete cds	96	450	450
2	6	6592	6167	emb 283335 SP28	S.pneumoniae dexB, cap1A, B, C, D, E, F, G, H, I, J, K) genes, dtDP-rhamnose biosynthesis genes and alia gene	98	426	426
3	11	9770	9147	emb 283335 SP28	S.pneumoniae dexB, cap1A, B, C, D, E, F, G, H, I, J, K) genes, dtDP-rhamnose biosynthesis genes and alia gene	94	624	624
3	12	10489	9671	emb 283335 SP28	S.pneumoniae dexB, cap1A, B, C, D, E, F, G, H, I, J, K) genes, dtDP-rhamnose biosynthesis genes and alia gene	91	819	819
3	13	11546	12019	gb U43526	Streptococcus pneumoniae neuraminidase B (tranB) gene, complete cds, and neuraminidase (nanA) gene, partial cds	99	474	474
3	14	12017	11375	gb U43526	Streptococcus pneumoniae neuraminidase B (tranB) gene, complete cds, and neuraminidase (nanA) gene, partial cds	99	1359	1359
3	15	13421	14338	gb U43526	Streptococcus pneumoniae neuraminidase B (tranB) gene, complete cds, and neuraminidase (nanA) gene, partial cds	99	918	918
3	16	14329	15171	gb U43526	Streptococcus pneumoniae neuraminidase B (tranB) gene, complete cds, and neuraminidase (nanA) gene, partial cds	99	843	843
3	17	15112	17282	gb U43526	Streptococcus pneumoniae neuraminidase B (tranB) gene, complete cds, and neuraminidase (nanA) gene, partial cds	99	2151	2151
3	18	17267	18397	gb U43526	Streptococcus pneumoniae neuraminidase B (tranB) gene, complete cds, and neuraminidase (nanA) gene, partial cds	99	1069	1131
4	1	46	1188	emb Y11463 SPDN	Streptococcus pneumoniae dnaG, rpoD, cpoA genes and ORF3 and ORF5	99	1143	1143
4	2	1198	2529	emb Y11463 SPDN	Streptococcus pneumoniae dnaG, rpoD, cpoA genes and ORF3 and ORF5	99	876	1332
5	7	11297	11473	gb U41735	Streptococcus pneumoniae peptide methionine sulfoxide reductase (msrA) and homoserine kinase homolog (thrB) genes, complete cds	82	175	177
6	7	7125	7364	emb 277726 SP15	S.pneumoniae DNA for insertion sequence IS1318 (1372 bp)	93	238	240
6	8	7322	7570	emb Z77725 SP15	S.pneumoniae DNA for insertion sequence IS1381 (966 bp)	95	160	249
6	9	7533	7985	emb 277725 SP15	S.pneumoniae DNA for insertion sequence IS1381 (966 bp)	99	453	453
6	23	20197	19733	emb 283335 SP28	S.pneumoniae dexB, cap1A, B, C, D, E, F, G, H, I, J, K) genes, dtDP-rhamnose biosynthesis genes and alia gene	96	465	465
7	10	8305	7682	emb 283335 SP28	S.pneumoniae dexB, cap1A, B, C, D, E, F, G, H, I, J, K) genes, dtDP-rhamnose biosynthesis genes and alia gene	95	624	624

TABLE 1
S. pneumoniae - Coding regions containing known sequences

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	Percent HSP ident	HSP at nt length	ORF nt length
7	11	9024	8206	emb Z81335 SP28	<i>S. pneumoniae</i> dexB, cap1A, B, C, D, E, F, G, H, I, J, K genes, dtDP-rhamnose biosynthesis genes and alIA gene	95	819	819
10	13	9304	8078	gb L29323	<i>Streptococcus pneumoniae</i> methyl transferase (mtr) gene cluster, complete cds	93	513	1227
11	2	548	919	emb Z79391 SOOR	<i>S. pneumoniae</i> yorf[A, B, C, D, E], ftsL, pbpX and rgrR genes	99	316	372
11	3	892	1980	emb Z79391 SOOR	<i>S. pneumoniae</i> yorf[A, B, C, D, E], ftsL, pbpX and rgrR genes	99	1089	1089
11	5	3040	3477	emb Z79391 SOOR	<i>S. pneumoniae</i> yorf[A, B, C, D, E], ftsL, pbpX and rgrR genes	99	259	438
11	6	3480	3247	emb Z79391 SOOR	<i>S. pneumoniae</i> yorf[A, B, C, D, E], ftsL, pbpX and rgrR genes	99	234	234
11	7	3601	4557	emb Z79391 SOOR	<i>S. pneumoniae</i> yorf[A, B, C, D, E], ftsL, pbpX and rgrR genes	98	557	957
11	8	4506	4866	emb Z79391 SOOR	<i>S. pneumoniae</i> yorf[A, B, C, D, E], ftsL, pbpX and rgrR genes	99	381	381
11	9	4884	7142	emb X16167 SPPB	<i>Streptococcus pneumoniae</i> pbpX gene for penicillin binding protein 2X	99	259	259
11	10	7132	8124	emb X16167 SPPB	<i>Streptococcus pneumoniae</i> pbpX gene for penicillin binding protein 2X	98	70	93
13	1	53	1126	gb M31296	<i>S. pneumoniae</i> recP gene, complete cds	99	437	1074
14	3	1837	2148	emb Z81335 SP28	<i>S. pneumoniae</i> dexB, cap1A, B, C, D, E, F, G, H, I, J, K genes, dtDP-rhamnose biosynthesis genes and alIA gene	87	96	312
14	4	2518	2108	gb M36180	<i>Streptococcus pneumoniae</i> transposase, (ccmA and ccmB) and SAICAR synthetase (purC) genes, complete cds	98	411	411
15	9	8942	8511	gb U09239	<i>Streptococcus pneumoniae</i> type 19F capsular polysaccharide biosynthesis operon, (cps19FABCDEFGHJKLMNO) genes, complete cds, and alIA gene, partial cds	89	340	432
17	7	3910	3458	emb Z777226 SP1S	<i>S. pneumoniae</i> DNA for insertion sequence IS1118 (1372 bp)	98	453	453
17	8	4304	3873	emb Z77727 SP1S	<i>S. pneumoniae</i> DNA for insertion sequence IS1118 (823 bp)	96	182	412
19	1	41	529	emb X94909 SP1G	<i>S. pneumoniae</i> iga gene	75	168	489
19	2	554	757	gb L07752	<i>Streptococcus pneumoniae</i> attachment site (attB), DNA sequence	99	167	204
19	3	946	1827	gb L07752	<i>Streptococcus pneumoniae</i> attachment site (attB), DNA sequence	94	100	882
20	1	937	182	gb U33315	<i>Streptococcus pneumoniae</i> orfL gene, partial cds, competence stimulating peptide precursor (comC), histidine protein kinase (comD) and response regulator (comE) genes, complete cds, tRNA-Arg and tRNA-Gln genes	99	756	756
20	2	2271	931	gb U33315	<i>Streptococcus pneumoniae</i> orfL gene, partial cds, competence stimulating peptide precursor (comC), histidine protein kinase (comD) and response regulator (comE) genes, complete cds, tRNA-Arg and tRNA-Gln genes	98	1341	1341

TABLE 1
S. pneumoniae - Coding regions containing known sequences

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	percent indent	HSP nt length	ORF nt length
20	3	3175	2684	gb U76218	Streptococcus pneumoniae competence stimulating peptide precursor ComC (ComC), histidine kinase homolog ComD (ComD), and response regulator homolog ComE (comE) genes, complete cds	99	492	492
20	4	3322	4527	gb AF000658	Streptococcus pneumoniae R801 tRNA-Arg gene, partial sequence, and putative serine protease (sphtra), SPSPQJ (spspqj), initiator protein (spdnna) and beta subunit of DNA Polymerase III (spdnan) genes, complete cds	99	1206	1206
20	5	4573	5343	gb AF000658	Streptococcus pneumoniae R801 tRNA-Arg gene, partial sequence, and putative serine protease (sphtra), SPSPQJ (spspqj), initiator protein (spdnna) and beta subunit of DNA Polymerase III (spdnan) genes, complete cds	99	771	771
20	6	5332	6917	gb AF000658	Streptococcus pneumoniae R801 tRNA-Arg gene, partial sequence, and putative serine protease (sphtra), SPSPQJ (spspqj), initiator protein (spdnna) and beta subunit of DNA Polymerase III (spdnan) genes, complete cds	99	1386	1386
20	7	6395	8212	gb AF000658	Streptococcus pneumoniae R801 tRNA-Arg gene, partial sequence, and putative serine protease (sphtra), SPSPQJ (spspqj), initiator protein (spdnna) and beta subunit of DNA Polymerase III (spdnan) genes, complete cds	99	1218	1218
20	8	8214	8471	gb AF000658	Streptococcus pneumoniae R801 tRNA-Arg gene, partial sequence, and putative serine protease (sphtra), SPSPQJ (spspqj), initiator protein (spdnna) and beta subunit of DNA Polymerase III (spdnan) genes, complete cds	98	258	258
20	9	8334	9670	gb AF000658	Streptococcus pneumoniae R801 tRNA-Arg gene, partial sequence, and putative serine protease (sphtra), SPSPQJ (spspqj), initiator protein (spdnna) and beta subunit of DNA Polymerase III (spdnan) genes, complete cds	99	134	1137
22	14	11887	12267	emb 277726 SP15	<i>S. pneumoniae</i> DNA for insertion sequence IS1118 (1372 bp)	99	226	381
22	15	12208	12256	emb 277727 SP15	<i>S. pneumoniae</i> DNA for insertion sequence IS1118 (1323 bp)	97	353	453
22	16	13165	12662	emb 277726 SP15	<i>S. pneumoniae</i> DNA for insertion sequence IS1118 (1372 bp)	98	504	504
22	23	18398	18910	emb 286112 SP28	<i>S. pneumoniae</i> genes encoding galacturonosyl transferase and transposase and insertion sequence IS1155	95	463	513
22	24	18829	19299	emb 286112 SP28	<i>S. pneumoniae</i> genes encoding galacturonosyl transferase and transposase and insertion sequence IS1155	99	443	471
23	1	5624	4203	emb X52474 SPPL	<i>S. pneumoniae</i> plv gene for pneumolysin	99	1422	1422
23	6	6063	5629	gb H17717	<i>S. pneumoniae</i> pneumolysin gene, complete cds	98	197	435
26	1	5500	2	emb X94999 SP1G	<i>S. pneumoniae</i> IgA gene	87	3487	5499
26	2	5823	5584	gb U47687	Streptococcus pneumoniae immunoglobulin A1 protease (iga) gene, complete cds	99	151	240
26	3	6878	5685	gb U47687	Streptococcus pneumoniae immunoglobulin A1 protease (iga) gene, complete cds	100	50	1194

TABLE 1
S. pneumoniae - Coding regions containing known sequences

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	Percent HSP ident	HSP nt length	ORF nt length
26	8	14498	14854	emb 283335 SP28	<i>S. pneumoniae</i> dexB, cap1A, B,C,D,E,F,G,H,I,J,K genes, dtDP-rhamnose biosynthesis genes and alia gene	99	338	357
26	9	14763	14924	emb 283335 SP28	<i>S. pneumoniae</i> dexB, cap1A, B,C,D,E,F,G,H,I,J,K genes, dtDP-rhamnose biosynthesis genes and alia gene	100	94	162
26	10	14922	15173	gb U04047	Streptococcus pneumoniae SS2 dextran glucosidase gene and insertion sequence IS1202 transposase gene, complete cds	97	242	252
28	1	80	505	emb 283335 SP28	<i>S. pneumoniae</i> dexB, cap1A, B,C,D,E,F,G,H,I,J,K genes, dtDP-rhamnose biosynthesis genes and alia gene	99	426	426
28	2	503	952	gb U04047	Streptococcus pneumoniae SS2 dextran glucosidase gene and insertion sequence IS1202 transposase gene, complete cds	97	450	450
28	3	780	1298	gb U04047	Streptococcus pneumoniae SS2 dextran glucosidase gene and insertion sequence IS1202 transposase gene, complete cds	96	181	519
34	1	207	1523	gb L08611	Streptococcus pneumoniae maltose/maltodextrin uptake (malX) and two maltodextrin permease (malC and malD) genes, complete cds	99	1317	1317
34	2	1477	2367	gb L08611	Streptococcus pneumoniae maltose/maltodextrin uptake (malX) and two maltodextrin permease (malC and malD) genes, complete cds	96	795	891
34	3	2533	3120	gb L21856	Streptococcus pneumoniae malA gene, complete cds; malR gene, complete cds	96	446	828
34	4	2790	2647	gb L21856	Streptococcus pneumoniae malA gene, complete cds; malR gene, complete cds	98	137	144
34	5	3118	4416	gb L21856	Streptococcus pneumoniae malA gene, complete cds; malR gene, complete cds	96	999	999
34	9	7764	7507	gb U41735	Streptococcus pneumoniae peptide methionine sulfoxide reductase (msrA) and homoserine kinase homolog (thrB) genes, complete cds	93	201	258
34	16	10562	10257	emb X53602 SP80	<i>S. pneumoniae</i> rnsA-Box	92	238	306
35	4	1176	1439	emb 283335 SP28	<i>S. pneumoniae</i> dexB, cap1A, B,C,D,E,F,G,H,I,J,K genes, dtDP-rhamnose biosynthesis genes and alia gene	87	248	264
35	5	1458	1961	gb U09239	Streptococcus pneumoniae type 19F capsular polysaccharide biosynthesis operon, (cps19fABCDEFGHijklmno) genes, complete cds, and alia gene, partial cds	98	264	504
35	17	16172	15477	emb X85787 SPCP	<i>S. pneumoniae</i> dexB, cps1A, cps14B, cps14C, cps14D, cps14E, cps14F, cps14G, cps14H, cps14I, cps14J, cps14L, tsa genes	97	696	696
35	18	16261	16170	emb 283335 SP28	<i>S. pneumoniae</i> dexB, cap1A, B,C,D,E,F,G,H,I,J,K genes, dtDP-rhamnose biosynthesis genes and alia gene	86	792	792
35	19	17620	16871	gb U09239	Streptococcus pneumoniae type 19F capsular polysaccharide biosynthesis operon, (cps19fABCDEFGHijklmno) genes, complete cds, and alia gene, partial cds	83	750	750

TABLE 1
S. pneumoniae - Coding regions containing known sequences

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	percent ident	HSP nt length	ORF nt length
35	20	19061	17664	emb X85787 SPCP	<i>S.pneumoniae</i> dxeB, cps1A, cps14B, cps14C, cps14D, cps14E, cps14F, cps14G, cps14H, cps14I, cps14J, cps14K, cps14L, tsa3 genes	94	1458	1458
36	19	18960	18352	gb U40786	streptococcus pneumoniae surface antigen A variant precursor (psaA) and 18 kDa protein genes, complete cds, and ORF1 gene, partial cds	99	609	609
36	20	1994	18366	gb U53509	streptococcus pneumoniae surface adhesin A precursor (psaA) gene, complete cds	99	969	969
37	1	2743	179	emb Z67739 SPPA	<i>S.pneumoniae</i> parC, parE and transposase genes and unknown orf	99	2565	2565
37	2	2985	2824	emb Z67739 SPPA	<i>S.pneumoniae</i> parC, parE and transposase genes and unknown orf	100	162	162
37	3	5014	3070	emb Z67739 SPPA	<i>S.pneumoniae</i> parC, parE and transposase genes and unknown orf	99	1965	1965
37	4	5134	5790	emb Z67739 SPPA	<i>S.pneumoniae</i> parC, parE and transposase genes and unknown orf	99	657	657
37	5	6171	5833	emb Z67739 SPPA	<i>S.pneumoniae</i> parC, parE and transposase genes and unknown orf	96	339	339
38	19	12269	11268	gb M28679	<i>S.pneumoniae</i> promoter region DNA	100	64	300
39	2	1256	2137	gb U41735	streptococcus pneumoniae peptide methionine sulfoxide reductase (msra) and homoserine kinase homolog (thrB) genes, complete cds	99	882	882
39	3	2405	3370	gb U41735	streptococcus pneumoniae peptide methionine sulfoxide reductase (msra) and homoserine kinase homolog (thrB) genes, complete cds	99	966	966
40	9	5253	7208	gb M29686	<i>S.pneumoniae</i> mismatch repair (hexB) gene, complete cds	99	1956	1956
41	1	3	1037	emb Z17307 SPRE	<i>S.pneumoniae</i> recA gene encoding RecA	99	1027	1035
41	2	1328	2713	emb Z24303 SPCR	streptococcus pneumoniae cin operon encoding the cinA, recA, dinF, lytA genes, and downstream sequences	99	1386	1386
41	3	3083	4045	gb M13812	<i>S.pneumoniae</i> autolysin (lytA) gene, complete cds	99	963	963
41	4	3272	3096	gb M13812	<i>S.pneumoniae</i> autolysin (lytA) gene, complete cds	100	177	177
41	5	3603	3860	gb M13812	<i>S.pneumoniae</i> autolysin (lytA) gene, complete cds	100	258	258
41	6	4755	5162	gb L36660	streptococcus pneumoniae ORF, complete cds	98	408	408
41	7	5270	5716	gb L36660	streptococcus pneumoniae ORF, complete cds	98	447	447
41	8	6112	6918	gb L36660	streptococcus pneumoniae ORF, complete cds	98	431	807
41	9	6916	7119	gb L36660	streptococcus pneumoniae ORF, complete cds	100	204	204
41	10	7082	7660	gb L36660	streptococcus pneumoniae ORF, complete cds	97	552	579
41	11	7680	7979	gb L36660	streptococcus pneumoniae ORF, complete cds	98	81	100
41	12	9169	8717	emb Z77727 SPIS	<i>S.pneumoniae</i> DNA for insertion sequence IS1118 (823 bp)	97	353	453

TABLE 1
S. pneumoniae - Coding regions containing known sequences

Contig		ORF	start	stop	match	match gene name	percent	HSP nt	ORF nt
ID	ID	ID	(nt)	(nt)	accession		ident	length	length
41	13	9533	9132	emb 277725 SPZ8	[S.pneumoniae DNA for insertion sequence IS1381 (1966 bp)		95	160	402
41	14	9669	9475	emb 282001 SPZ8	[S.pneumoniae pcpA gene and open reading frames		100	189	195
44	5	7190	7555	emb 282001 SPZ8	[S.pneumoniae pcpA gene and open reading frames		99	166	166
44	6	8059	7607	emb 277726 SPZ8	[S.pneumoniae DNA for insertion sequence IS1318 (11372 bp)		97	453	453
44	7	8423	8022	emb 277725 SPZ8	[S.pneumoniae DNA for insertion sequence IS1381 (1966 bp)		95	160	402
44	8	8559	8365	emb 282001 SPZ8	[S.pneumoniae pcpA gene and open reading frames		100	189	195
48	9	6480	4687	gb L39074	[Streptococcus pneumoniae pyruvate oxidase (spxB) gene, complete cds		99	1794	1794
49	2	231	2603	gb L20561	[Streptococcus pneumoniae Exp7 gene, partial cds		100	216	2373
53	6	2407	2156	gb U04047	[Streptococcus pneumoniae Sz2 dextran glucosidase gene and insertion sequence IS1202 transpose gene, complete cds		97	242	252
53	7	2566	2405	emb 283335 SPZ8	[S.pneumoniae dexB, capIa, p, c, d, e, f, g, h, i, j, k] genes, drDP-rhamnose biosynthesis genes and aiaA gene		100	94	162
53	8	2831	2475	emb 283335 SPZ8	[S.pneumoniae dexB; capIa, p, c, d, e, f, g, h, i, j, k] genes, drDP-rhamnose biosynthesis genes and aiaA gene		99	138	357
54	13	12409	11105	emb 283335 SPZ8	[S.pneumoniae dexB, capIa, p, c, d, e, f, g, h, i, j, k] genes, drDP-rhamnose biosynthesis genes and aiaA gene		67	591	1305
55	22	120488	119449	emb 1284379 HS28	[S.pneumoniae dfr gene (isolate 92)]		99	540	540
61	11	11864	9900	emb 216082 PNAL	[Streptococcus pneumoniae aibA gene		98	1965	1965
63	1	3	239	gb M18729	[S.pneumoniae mismatch repair protein (hexA) gene, complete cds		100	237	237
63	2	233	2611	gb M18729	[S.pneumoniae mismatch repair protein (hexA) gene, complete cds		99	230	239
63	3	2557	2823	gb M18729	[S.pneumoniae mismatch repair protein (hexA) gene, complete cds		99	266	267
63	4	2958	4664	gb M18729	[S.pneumoniae mismatch repair protein (hexA) gene, complete cds		95	69	1707
67	6	3770	3399	gb L220670	[Streptococcus pneumoniae hyaluronidase gene, complete cds		96	372	372
67	7	7161	4171	gb L220670	[Streptococcus pneumoniae hyaluronidase gene, complete cds		99	2938	2991
70	1	1	702	gb M14340	[S.pneumoniae DpnI gene region encoding dpmC and dpmB, complete cds		100	693	702
70	2	678	1160	gb M14340	[S.pneumoniae DpnI gene region encoding dpmC and dpmB, complete cds		100	483	483
70	3	2490	1210	gb M14339	[S.pneumoniae DpnII gene region encoding dpmN, dpmA, dpmB, complete cds		98	462	1281
70	7	4230	4424	gb J04234	[S.pneumoniae exodeoxyribonuclease (exoA) gene, complete cds		99	147	195
70	8	5197	4316	gb J04234	[S.pneumoniae exodeoxyribonuclease (exoA) gene, complete cds		99	881	882

TABLE 1
S. pneumoniae - Coding regions containing known sequences:

Contig	ORF ID	start (nt)	stop (nt)	match accession	match gene name	percent ident	HSP nt length	ORF nt length
70	113	8108	9874	gb L20562	Streptococcus pneumoniae Exp8 gene, partial cds	93	234	1767
71	22	127964	28341	emb X63602 SP0	S.pneumoniae msaA-Box	93	233	378
72	5	4697	3552	emb 226850 SPAT	S.pneumoniae (R222) genes for ATPase a subunit, ATPase b subunit and ATPase c subunit	97	102	1056
73	1	471	133	emb X63602 SP0	S.pneumoniae msaA-Box	91	193	339
73	3	3638	977	gb J04479	S.pneumoniae DNA polymerase I (polA) gene, complete cds	99	2682	2682
73	8	4884	5379	gb M36180	Streptococcus pneumoniae transposase, (comA and comB) and SaICAR synthetase	98	318	516
77	3	2652	1999	emb Z83335 SP28	S.pneumoniae dexB, capI[A,B,C,D,E,F,G,H,I,J,K] genes, dtDP-rihamnose biosynthesis genes and alia gene	95	624	624
77	4	3341	2523	emb Z83335 SP28	S.pneumoniae dexB, capI[A,B,C,D,E,F,G,H,I,J,K] genes, dtDP-rihamnose biosynthesis genes and alia gene	91	819	819
78	1	341	3	emb X77249 SPR6	S.pneumoniae (R6) ciaR/ciaH genes	99	139	139
78	2	1055	325	emb X77249 SPR6	S.pneumoniae (R6) ciaR/ciaH genes	99	771	771
82	10	11436	10816	gb U90721	Streptococcus pneumoniae signal peptidase I (spI) gene, complete cds	97	621	621
82	11	12402	11434	gb U93576	Streptococcus pneumoniae ribonuclease HII (rnhb) gene, complete cds	98	953	969
82	12	12381	112704	gb U93576	Streptococcus pneumoniae ribonuclease HII (rnhb) gene, complete cds	100	51	324
83	8	3212	3550	emb 277727 SP15	S.pneumoniae DNA for insertion sequence IS1318 (1823 bp)	97	290	339
83	10	4662	6851	gb M36180	Streptococcus pneumoniae transposase, (comA and comB) and SaICAR synthetase	99	2190	2190
83	11	6849	8213	gb M36180	Streptococcus pneumoniae transposase, (purC) genes, complete cds	99	1365	1365
83	12	8226	9090	gb M36180	Streptococcus pneumoniae transposase, (comA and comB) and SaICAR synthetase	99	855	855
83	13	9233	13017	gb L15190	Streptococcus pneumoniae SaICAR synthetase (purC) gene, complete cds	100	107	1735
83	23	22147	23313	gb L36923	Streptococcus pneumoniae beta-N-acetylhexosaminidase (stII) gene, complete cds	98	218	1167
83	24	23268	23450	gb L36923	Streptococcus pneumoniae beta-N-acetylhexosaminidase (stII) gene, complete cds	98	172	183
83	25	27527	23505	gb L36923	Streptococcus pneumoniae beta-N-acetylhexosaminidase (stII) gene, complete cds	99	3826	4023

TABLE 1 S. pneumoniae - Coding regions containing known sequences

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	percent ident	HSP nt length	ORF nt length
83	26	28872	27771	[gb]L36921]	streptococcus pneumoniae beta-N-acetylhexosaminidase (strH) gene, complete cds	99	416	702
84	4	4554	6173	[emb]283335 SP28	S.pneumoniae dexB, cap1A, B, C, D, E, F, G, H, I, J, K1 genes, dTDP-β-hamnose biosynthesis genes and alia gene	98	697	1620
87	6	5951	5316	[emb]277725 SP28	S.pneumoniae DNA for insertion sequence IS1381 (966 bp)	96	439	636
88	5	2957	3511	[gb]M16180]	streptococcus pneumoniae transposase, (conA and comB) and SAICAR synthetase genes, complete cds	94	555	555
88	6	3466	4269	[gb]M16180]	streptococcus pneumoniae transposase, (conA and comB) and SAICAR synthetase genes, complete cds	94	804	804
89	13	9878	10093	[gb]M16180]	streptococcus pneumoniae transposase, (conA and comB) and SAICAR synthetase genes, complete cds	97	211	216
89	14	10062	10412	[emb]283335 SP28	S.pneumoniae dexB, cap1A, B, C, D, E, F, G, H, I, J, K1 genes, dTDP-β-hamnose biosynthesis genes and alia gene	97	315	351
93	10	5303	4941	[emb]X63602 SPB0	S.pneumoniae mmsA-Box	89	237	363
97	4	1708	1520	[gb]U41735]	streptococcus pneumoniae peptide methionine sulfoxide reductase (msrA) and homoserine kinase homolog (chvB) genes, complete cds	91	140	189
99	1	89	700	[emb]283335 SP28	S.pneumoniae dexB, cap1A, B, C, D, E, F, G, H, I, J, K1 genes, dTDP-β-hamnose biosynthesis genes and alia gene	93	592	612
99	2	1773	775	[emb]X17337 SPAM	streptococcus pneumoniae aml locus conferring aminopterin resistance	99	998	999
99	3	2794	1712	[emb]X17337 SPAM	streptococcus pneumoniae aml locus conferring aminopterin resistance	99	1083	1083
99	4	3732	2788	[emb]X17337 SPAM	streptococcus pneumoniae aml locus conferring aminopterin resistance	100	945	945
99	5	5249	3714	[emb]X17337 SPAM	streptococcus pneumoniae aml locus conferring aminopterin resistance	100	1536	1536
99	6	7262	5277	[emb]X17337 SPAM	streptococcus pneumoniae aml locus conferring aminopterin resistance	99	1986	1986
101	1	216	1538	[emb]X54225 SPEN	S.pneumoniae epuA and endA genes for 7 kDa protein and membrane endonuclease	99	146	1323
101	2	1492	1719	[emb]X54225 SPEN	S.pneumoniae epuA and endA genes for 7 kDa protein and membrane endonuclease	100	162	162
101	4	1701	2582	[emb]X54225 SPEN	S.pneumoniae epuA and endA genes for 7 kDa protein and membrane endonuclease	100	882	882
103	7	5556	5041	[emb]295914 SP29	streptococcus pneumoniae sodA gene	100	396	516
104	2	1347	1556	[emb]27727 SPIS	S.pneumoniae DNA for insertion sequence IS1318 (1823 bp)	83	206	210

TABLE 1
S. pneumoniae - Coding regions containing known sequences

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	HSP nt ident	ORF nt length
105	5	5381	5028	emb 267739 SPPA	<i>S. pneumoniae</i> parC, parE and transposase genes and unknown orf	98	353
105	6	6089	5379	emb 267739 SPPA	<i>S. pneumoniae</i> parC, parE and transposase genes and unknown orf	98	84
107	4	2785	1880	emb X16022 SPPE	<i>S. pneumoniae</i> parA gene	98	72
107	5	2913	4988	emb X16022 SPPE	<i>S. pneumoniae</i> parA gene	99	1692
107	6	4981	5595	emb X11136 SPPE	streptococcus pneumoniae parA gene for penicillin binding protein 2B (penicillin resistant strain) lacking N-term.	91	107
108	9	9068	8718	emb 267739 SPPA	<i>S. pneumoniae</i> parC, parE and transposase genes and unknown orf	95	142
108	12	11308	10922	emb 267739 SPPA	<i>S. pneumoniae</i> parC, parE and transposase genes and unknown orf	99	199
109	3	2768	2241	emb 277725 SPIS	<i>S. pneumoniae</i> DNA for insertion sequence IS1361 (966 bp)	96	61
109	4	2688	2855	emb 277726 SPIS	<i>S. pneumoniae</i> DNA for insertion sequence IS1318 (11372 bp)	96	148
109	5	2862	3269	emb 277727 SPIS	<i>S. pneumoniae</i> DNA for insertion sequence IS1318 (823 bp)	97	353
109	6	5120	3584	gb M18729	<i>S. pneumoniae</i> mismatch repair protein (hexA) gene, complete cds	100	371
113	1	431	3	gb M16180	streptococcus pneumoniae transposase, (comA and comB) and SMICAR synthetase (purC) genes, complete cds	95	429
113	10	9788	8532	emb X99400 SPDA	<i>S. pneumoniae</i> dacA gene and ORF (purC) genes, complete cds	99	1257
113	11	9870	10985	emb X99400 SPDA	<i>S. pneumoniae</i> dacA gene and ORF (purC) genes, complete cds	99	1116
114	3	2530	2030	gb M16180	streptococcus pneumoniae transposase, (comA and comB) and SMICAR synthetase (purC) genes, complete cds	95	481
115	11	11303	10932	gb U04047	streptococcus pneumoniae SSZ dextran glucosidase gene and insertion sequence IS1202 transposase gene, complete cds	97	372
117	1	897	3102	emb X72967 SPNA	<i>S. pneumoniae</i> nana gene	99	2402
117	2	3277	3831	emb X72967 SPNA	<i>S. pneumoniae</i> nana gene	99	237
117	3	4327	3899	gb M16180	streptococcus pneumoniae transposase, (purC) genes, complete cds	98	429
121	2	1169	1941	gb U272720	streptococcus pneumoniae heat shock protein 70 (dnak) gene, complete cds and DnaJ (dnaj) gene, partial cds	99	202
121	3	2412	4253	gb U272720	streptococcus pneumoniae heat shock protein 70 (dnak) gene, complete cds and DnaJ (dnaj) gene, partial cds	99	1842
122	8	5066	5587	gb U04047	streptococcus pneumoniae SSZ dextran glucosidase gene and insertion sequence IS1202 transposase gene, complete cds	64	451

TABLE 1
S. pneumoniae - Coding regions containing known sequences

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	Percent HSP ident	HSP nt length	ORF nt length
125	1	1811	189	gb M36180	streptococcus pneumoniae transposase, (ccmA and comB) and SAICAR synthetase (purC) genes, complete cds	92	99	1623
128	15	12496	11204	emb 283335 SP28	<i>S. pneumoniae</i> dexB, capI [A,B,C,D,E,F,G,H,I,J,K] genes, dtDP-phamnose biosynthesis genes and alia gene	91	705	1293
134	1	1	492	emb Y10818 SPY1	<i>S. pneumoniae</i> spsA gene	99	203	492
134	2	556	2652	gb AF019904	streptococcus pneumoniae choline binding protein A (cbpA) gene, partial cds	86	685	2097
134	3	1160	817	emb Y10818 SPY1	<i>S. pneumoniae</i> spsA gene	86	324	324
134	4	3932	2882	gb AF019904	streptococcus pneumoniae choline binding protein A (cbpA) gene, partial cds	98	215	1071
134	8	7392	9848	gb U12567	streptococcus pneumoniae PI3 glycerol-3-phosphate dehydrogenase (glpD) gene, partial cds, and glycerol uptake facilitator (glpF) and ORF3 genes, complete cds	99	285	1857
134	9	9846	10622	gb U12567	streptococcus pneumoniae PI3 glycerol-3-phosphate dehydrogenase (glpD) gene, partial cds, and glycerol uptake facilitator (glpF) and ORF3 genes, complete cds	99	570	777
134	10	110305	11122	gb U12567	streptococcus pneumoniae PI3 glycerol-3-phosphate dehydrogenase (glpD) gene, partial cds, and glycerol uptake facilitator (glpF) and ORF3 genes, complete cds	100	318	318
137	13	7970	8443	gb U002339	streptococcus pneumoniae type 19F capsular polysaccharide biosynthesis operon, (cps19FABCDEFGHJKLMNO) genes, complete cds, and alia gene, partial cds	90	420	474
137	14	8590	8775	emb 283335 SP28	<i>S. pneumoniae</i> dexB, capI [A,B,C,D,E,F,G,H,I,J,K] genes, dtDP-phamnose biosynthesis genes and alia gene	94	174	186
137	15	8773	8967	emb 283335 SP28	<i>S. pneumoniae</i> dexB, capI [A,B,C,D,E,F,G,H,I,J,K] genes, dtDP-phamnose biosynthesis genes and alia gene	98	195	195
137	16	9223	9687	emb 277726 SP7S	<i>S. pneumoniae</i> DNA for insertion sequence IS1318 (1372 bp)	96	446	465
137	17	9641	10051	emb 277727 SP7S	<i>S. pneumoniae</i> DNA for insertion sequence IS1318 (823 bp)	96	293	411
139	10	12398	12702	emb X63602 SPBO	<i>S. pneumoniae</i> mnsA-Box	90	234	297
141	8	7805	8938	emb 249988 SPMM	streptococcus pneumoniae mnsA gene	100	76	936
141	9	8936	10972	emb 249988 SPMM	streptococcus pneumoniae mnsA gene	99	338	1134
141	10	11472	12467	emb 249988 SPMM	streptococcus pneumoniae mnsA gene	99	2037	2037
142	2	257	814	gb HE0215	streptococcus pneumoniae uvs402 protein gene, complete cds	98	174	558
142	3	787	957	gb HE0215	streptococcus pneumoniae uvs402 protein gene, complete cds	100	142	171
142	4	980	3022	gb HE0215	streptococcus pneumoniae uvs402 protein gene, complete cds	95	1997	2043

TABLE 1
S. pneumoniae - Coding regions containing known sequences

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	percent	HSP nt	ORF nt
						ident	length	length
142	5	3020	3595	gb M80215	Streptococcus pneumoniae uvs02 protein gene, complete cds	100	153	576
145	1	1	219	emb 25135 SPAL	<i>S. pneumoniae</i> alIA gene for amA-like gene A	97	185	219
145	2	171	1994	gb L20556	Streptococcus pneumoniae PlpA gene, partial cds	99	1811	1824
145	3	2387	7599	emb 247210 SPDE	<i>S. pneumoniae</i> dexB, cap3A, cap3B and cap3C genes and orfs	99	1052	5313
145	4	9334	7766	gb M90527	Streptococcus pneumoniae penicillin-binding protein (ponA) gene, complete cds	99	2169	2169
145	5	1088	9922	gb M90527	Streptococcus pneumoniae penicillin-binding protein (ponA) gene, complete cds	99	512	567
146	1	159	4	emb 282002 SPZB	<i>S. pneumoniae</i> pcpB and pcpC genes	98	156	156
146	2	344	90	emb 282002 SPZB	<i>S. pneumoniae</i> pcpB and pcpC genes	98	255	255
146	16	11195	10794	emb 282002 SPZB	<i>S. pneumoniae</i> pcpB and pcpC genes	85	276	1002
147	11	10378	10202	emb 221702 SPUN	<i>S. pneumoniae</i> ung gene and mutX genes encoding uracil-DNA glycosylase and 8-oxodGTP nucleoside triphosphatase	98	477	477
147	12	111338	10676	emb 221702 SPUN	<i>S. pneumoniae</i> ung gene and mutX genes encoding uracil-DNA glycosylase and 8-oxodGTP nucleoside triphosphatase	99	663	663
148	12	9009	8815	gb U41735	Streptococcus pneumoniae Peptidyl methionine sulfoxide reductase (msrA) and homoserine kinase homolog (thrB) genes, complete cds	90	180	195
156	4	1154	1402	emb X6362 SPBO	<i>S. pneumoniae</i> umsA-Box	94	185	249
159	13	9048	8521	gb M56180	Streptococcus pneumoniae transposase, (conA and comB) and SICAR synthetase (purC) genes, complete cds	98	526	528
160	1	1	147	emb 226831 SPAT	<i>S. pneumoniae</i> (R6) genes for ATPase a subunit, ATPase b subunit and ATPase c subunit	100	142	147
160	2	179	898	emb 226831 SPAT	<i>S. pneumoniae</i> (R6) genes for ATPase a subunit, ATPase b subunit and ATPase c subunit	99	720	720
160	3	906	1406	emb 226830 SPAT	<i>S. pneumoniae</i> (M22) genes for ATPase a subunit, ATPase b subunit and ATPase c subunit	95	501	501
160	4	1373	1942	emb 226830 SPAT	<i>S. pneumoniae</i> (M22) genes for ATPase a subunit, ATPase b subunit and ATPase c subunit	87	306	570
161	1	1	984	emb X7249 SPR6	<i>S. pneumoniae</i> (R6) clpR/clpH genes	99	984	984
161	7	6910	7497	emb X83917 SPGY	<i>S. pneumoniae</i> orf1gyrB and gyrB gene encoding DNA gyrase B subunit	99	417	588
161	8	7443	9386	emb X83917 SPGY	<i>S. pneumoniae</i> orf1gyrB and gyrB gene encoding DNA gyrase B subunit	98	1912	1944
163	1	2	2155	gb L20559	Streptococcus pneumoniae ExpS gene, partial cds	98	327	2154

TABLE I
S. pneumoniae - Coding regions containing known sequences

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	percent HSP ident	HSP nt length	ORF nt length
165	1	32	1618	gb J01796	<i>S. pneumoniae</i> malX and malM genes encoding membrane protein and amylo-ltase, complete cds, and malP gene encoding phosphorlyase	99	1587	1587
165	2	1608	3902	gb J01796	<i>S. pneumoniae</i> malX and malM genes encoding membrane protein and amylo-ltase, complete cds, and malP gene encoding phosphorlyase	100	280	2295
166	1	378	4	emb Y11463 SPDN	<i>Streptococcus pneumoniae</i> dnaG, rpoD, cpol genes and ORF3 and ORF5	100	375	375
166	2	1507	320	emb Y11463 SPDN	<i>Streptococcus pneumoniae</i> dnaG, rpoD, cpol genes and ORF3 and ORF5	99	1188	1188
166	3	3240	1432	emb Y11463 SPDN	<i>Streptococcus pneumoniae</i> dnaG, rpoB, cpol genes and ORF3 and ORF5	99	563	1009
167	1	1077	328	emb Z71552 SPAD	<i>Streptococcus pneumoniae</i> adCCBA operon	94	155	750
167	2	1844	999	emb Z71552 SPAD	<i>Streptococcus pneumoniae</i> adCCBA operon	98	405	846
167	3	2714	1842	emb Z71552 SPAD	<i>Streptococcus pneumoniae</i> adCCBA operon	97	604	873
167	4	3399	261	emb Z71552 SPAD	<i>Streptococcus pneumoniae</i> adCCBA operon	99	703	759
168	1	1	2259	gb L2058	<i>Streptococcus pneumoniae</i> Exp4 gene, partial cds	99	282	2259
170	10	7338	7685	emb Z77726 SPIS	<i>S. pneumoniae</i> DNA for insertion sequence IS1318 (1372 bp)	95	315	148
172	6	2462	4981	gb U47625	<i>Streptococcus pneumoniae</i> formate acetyltransferase (exp72) gene, partial cds	97	365	2520
175	1	373	20	gb M36180	<i>Streptococcus pneumoniae</i> transposase, lcomA and comB and SAICAR synthetase (purC) genes, complete cds	89	153	354
175	4	183	3621	emb Z47210 SPDE	<i>S. pneumoniae</i> dpxB, cap1B and cap1C genes and orfs	95	89	1779
176	5	3984	280	emb Z67739 SPPA	<i>S. pneumoniae</i> parC, parE and transposase genes and unknown orf	100	573	1005
178	1	3	425	emb Z67739 SPPA	<i>S. pneumoniae</i> parC, parE and transposase genes and unknown orf	95	423	423
179	1	426	70	emb Z83335 SP28	<i>S. pneumoniae</i> dpxB, cap1(A,B,C,D,E,F,G,H,I,J,K) genes, dpxB-rhamnose biosynthesis genes and alIA gene	99	338	357
180	3	3084	1855	emb X55718 SP0Y	<i>S. pneumoniae</i> gyra gene	99	381	1230
186	1	714	4	emb Z79691 SOOR	<i>S. pneumoniae</i> yorf1A, B,C,D,E, ftsL, pbpX and regR genes	98	59	711
186	2	2254	608	emb Z79691 SOOR	<i>S. pneumoniae</i> yorf1A, B,C,D,E, ftsL, pbpX and regR genes	98	315	1647
186	3	707	880	emb Z79691 SOOR	<i>S. pneumoniae</i> yorf1A, B,C,D,E, ftsL, pbpX and regR genes	98	174	174
189	1	2	259	gb U72720	<i>Streptococcus pneumoniae</i> heat shock protein 70 (dnak) gene, complete cds and DnaJ (dnaj) gene, partial cds	99	258	258
189	2	600	385	gb U72720	<i>Streptococcus pneumoniae</i> heat shock protein 70 (dnak) gene, complete cds and DnaJ (dnaj) gene, partial cds	98	244	246

TABLE 1
S. pneumoniae - Coding regions containing known sequences

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	Percent ident	HSP nt length	ORF nt length
189	3	1018	851	gb U72720	streptococcus pneumoniae heat shock protein 70 (dnak) gene, partial cds	99	168	168
189	4	1012	2154	gb U72720	streptococcus pneumoniae heat shock protein 70 (dnak) gene, partial cds and DnaJ (dnaj) gene, partial cds	99	1062	1143
191	9	7829	7524	emb X63602 SPB0	<i>S. pneumoniae</i> mmsA-Box	95	234	306
194	1	1	729	gb M36180	streptococcus pneumoniae transposase, (conA and comB) and SAICAR synthetase (purC) genes, complete cds	91	728	729
199	2	1117	881	emb Z83335 SP28	<i>S. pneumoniae</i> daxB, caplA,B,C,D,E,F,G,H,I,J,K genes, dtDP-rhamnose biosynthesis genes and alIA gene	96	211	237
199	4	1499	1762	emb Z83335 SP28	<i>S. pneumoniae</i> daxB, caplA,B,C,D,E,F,G,H,I,J,K genes, dtDP-rhamnose biosynthesis genes and alIA gene	89	248	264
199	5	1781	2280	emb Z83335 SP28	<i>S. pneumoniae</i> daxB, caplA,B,C,D,E,F,G,H,I,J,K genes, dtDP-rhamnose biosynthesis genes and alIA gene	98	504	504
203	1	1977	337	gb L20531	streptococcus pneumoniae Exp9 gene, partial cds	99	342	1641
204	1	1145	3	gb J3.6131	streptococcus pneumoniae exp10 gene, complete cds, recA gene, 5' end	99	1143	1143
208	1	59	2296	gb U89711	streptococcus pneumoniae pneumococcal surface protein A (pspa) gene, complete cds	90	471	2238
213	3	2455	2123	emb Z83335 SP28	<i>S. pneumoniae</i> daxB, caplA,B,C,D,E,F,G,H,I,J,K genes, dtDP-rhamnose biosynthesis genes and alIA gene	96	332	333
216	1	368	12	emb Z83335 SP28	<i>S. pneumoniae</i> daxB, caplA,B,C,D,E,F,G,H,I,J,K genes, dtDP-rhamnose biosynthesis genes and alIA gene	99	338	357
216	3	2650	2327	gb M28678	<i>S. pneumoniae</i> promoter sequence DNA	98	86	124
222	1	417	4	emb Z83335 SP28	<i>S. pneumoniae</i> daxB, caplA,B,C,D,E,F,G,H,I,J,K genes, dtDP-rhamnose biosynthesis genes and alIA gene	94	414	414
227	1	5266	4228	emb AJ003326 SP	streptococcus pneumoniae 1dh gene	99	1029	1029
239	1	1	804	gb M31296	<i>S. pneumoniae</i> recP gene, complete cds	95	484	804
247	3	1625	1807	gb M36180	streptococcus pneumoniae transposase, (conA and comB) and SAICAR synthetase (purC) genes, complete cds	94	178	183
249	3	921	1364	emb Z83335 SP28	<i>S. pneumoniae</i> daxB, caplA,B,C,D,E,F,G,H,I,J,K genes, dtDP-rhamnose biosynthesis genes and alIA gene	94	443	444
253	1	362	3	gb M36180	streptococcus pneumoniae transposase, (conA and comB) and SAICAR synthetase (purC) genes, complete cds	99	360	360
253	5	1218	2050	emb Z83335 SP28	<i>S. pneumoniae</i> daxB, caplA,B,C,D,E,F,G,H,I,J,K genes, dtDP-rhamnose biosynthesis genes and alIA gene	95	420	813

TABLE 1 *S. pneumoniae* - Coding regions containing known sequences

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	percent HSP nt ident	HSP nt length	ORF nt length
253	6	2069	2572	emb 28335 SP28	<i>S. pneumoniae</i> dexB, cap1A, B, C, D, E, F, G, H, I, J, K1 genes, dTDP-rhamnose biosynthesis genes and alia gene	97	504	504
255	1	3	800	emb 28202 SP28	<i>S. pneumoniae</i> pcP8 and pcPc genes	97	531	798
255	2	798	1841	emb 28202 SP28	<i>S. pneumoniae</i> pcP8 and pcPc genes	97	672	1044
255	3	2493	1969	emb 267739 SPPA	<i>S. pneumoniae</i> parC, parE and transposase genes and unknown orf	92	435	525
257	2	985	770	emb X17337 SPAM	<i>Streptococcus pneumoniae</i> ami locus conferring aminopterin resistance	96	117	216
257	3	1245	907	gb M36180	Streptococcus pneumoniae transposase, (cmaA and cmaB) and SAICAR synthetase (pURC) genes, complete cds	97	339	339
267	2	495	1208	gb U16156	Streptococcus pneumoniae dihydropteroate synthase (sulA), dihydrofolate synthetase (sulB), guanosine triphosphate cyclohydrolase (sulC), aldoles-pyrophosphokinase (sulD) genes, complete cds	95	84	714
267	3	1291	2277	gb U16156	Streptococcus pneumoniae dihydropteroate synthase (sulA), dihydrofolate synthetase (sulB), guanosine triphosphate cyclohydrolase (sulC), aldoles-pyrophosphokinase (sulD) genes, complete cds	97	755	987
267	4	2261	3601	gb U16156	Streptococcus pneumoniae dihydropteroate synthase (sulA), dihydrofolate synthetase (sulB), guanosine triphosphate cyclohydrolase (sulC), aldoles-pyrophosphokinase (sulD) genes, complete cds	98	1341	1341
267	5	3561	4136	gb U16156	Streptococcus pneumoniae dihydropteroate synthase (sulA), dihydrofolate synthetase (sulB), guanosine triphosphate cyclohydrolase (sulC), aldoles-pyrophosphokinase (sulD) genes, complete cds	99	576	576
267	6	4164	4949	gb U16156	Streptococcus pneumoniae dihydropteroate synthase (sulA), dihydrofolate synthetase (sulB), guanosine triphosphate cyclohydrolase (sulC), aldoles-pyrophosphokinase (sulD) genes, complete cds	99	748	786
267	7	5544	5140	gb U16156	Streptococcus pneumoniae dihydropteroate synthase (sulA), dihydrofolate synthetase (sulB), guanosine triphosphate cyclohydrolase (sulC), aldoles-pyrophosphokinase (sulD) genes, complete cds	100	186	405
268	4	1793	1990	emb X33602 SPBO	<i>S. pneumoniae</i> rnsA-Box	89	194	198
271	1	562	104	gb M28686	<i>S. pneumoniae</i> mismatch repair (hexB) gene, complete cds	93	160	459
291	1	75	524	gb U0047	Streptococcus pneumoniae SS2 dextran glucosidase gene and insertion sequence IS1202 transposase gene, complete cds	96	450	450
291	2	1001	525	emb 283335 SP28	<i>S. pneumoniae</i> dexB, cap1A, B, C, D, E, F, G, H, I, J, K1 genes, dTDP-rhamnose biosynthesis genes and alia gene	87	205	477
291	3	807	559	emb 283335 SP28	<i>S. pneumoniae</i> dexB, cap1A, B, C, D, E, F, G, H, I, J, K1 genes, dTDP-rhamnose biosynthesis genes and alia gene	90	170	249
291	4	1374	1099	gb M426180	<i>Streptococcus pneumoniae</i> transposase, (cmaA and cmaB) and SAICAR synthetase (pURC) genes, complete cds	85	264	276

TABLE 1
S. pneumoniae - Coding regions containing known sequences

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	Percent ident	RSP nt length	ORF nt length
293	1	3	1673	emb 267710 SP04	<i>S. pneumoniae</i> gyrB gene and unknown orf	98	553	1671
296	1	1434	151	emb 247210 SP06	<i>S. pneumoniae</i> dexB, cap3A, cap3B and cop3C genes and orfs	99	430	1284
317	1	157	510	emb 267739 SP0A	<i>S. pneumoniae</i> parC, parE and transposase genes and unknown orf	89	353	354
325	2	1237	485	emb 283315 SP28	<i>S. pneumoniae</i> dexB, cap1A,B,C,D,E,F,G,H,I,J,K genes, dtDP-rhamnose biosynthesis genes and alia gene	91	239	753
326	1	1	462	emb 282001 SP28	<i>S. pneumoniae</i> pcpA gene and open reading frames	100	233	462
327	1	603	64	emb 283315 SP28	<i>S. pneumoniae</i> dexB, cap1A,B,C,D,E,F,G,H,I,J,K genes, dtDP-rhamnose biosynthesis genes and alia gene	94	89	540
334	1	153	545	gb U41715	Streptococcus pneumoniae peptide methionine sulfoxide reductase (msrA) and homoserine kinase homolog (thrB) genes, complete cds	87	91	393
336	1	308	93	emb Z268501 SPAT	<i>S. pneumoniae</i> (R222) genes for ATPase a subunit, ATPase b subunit and ATPase c subunit	97	102	216
360	1	1	519	emb Z67739 SP0A	<i>S. pneumoniae</i> parC, parE and transposase genes and unknown orf	95	435	519
360	4	1598	1960	emb Z83335 SP28	<i>S. pneumoniae</i> dexB, cap1A,B,C,D,E,F,G,H,I,J,K genes, dtDP-rhamnose biosynthesis genes and alia gene	94	353	363
362	1	673	2	emb 283315 SP28	<i>S. pneumoniae</i> dexB, cap1A,B,C,D,E,F,G,H,I,J,K genes, dtDP-rhamnose biosynthesis genes and alia gene	95	63	672
362	2	1168	728	gb U04047	Streptococcus pneumoniae SS2 dextran glucosidase gene and insertion sequence IS1202 transposase gene, complete cds	96	441	441
384	1	347	111	emb X85787 SPCP	<i>S. pneumoniae</i> dexB, cps14A, cps14B, cps14C, cps14D, cps14E, cps14F, cps14G, cps14H, cps14I, cps14J, cps14K, cps14L, tasa genes	94	54	237

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
228	2	1760	1942	pir F6066 F606	translation elongation factor Tu - <i>Streptococcus oralis</i>	100	100	183
319	1	2	205	gi 1984927	neoycin phosphotransferase [Cloning vector pBSL99]	100	100	204
260	1	2	1138	pir F6066 F606	translation elongation factor Tu - <i>Streptococcus oralis</i>	99	98	1137
25	2	486	1394	gi 1157495	hypothetical [Haemophilus influenzae]	98	96	909
94	2	685	1002	gi 310627	phosphoenolpyruvate:sugar phosphotransferase system HPr [Streptococcus mutans]	98	93	318
312	1	190	2	gi 1347999	ATP-dependent protease proteolytic subunit [Streptococcus salivarius]	98	95	189
329	1	1	807	gi 1924848	inosine monophosphate dehydrogenase [Streptococcus pyogenes]	98	94	807
336	2	290	589	gi 987050	lacZ gene product [unidentified cloning vector]	98	98	300
181	9	5948	7366	gi 153755	phospho-beta-D-galactosidase [EC 3.2.1.85] [Lactococcus lactis cremoris]	97	94	1419
312	2	1044	361	gi 1347998	uracil phosphotriboyltransferase [Streptococcus salivarius]	97	88	684
32	8	6575	7486	sp P37214 ERA_S	GTP-BINDING PROTEIN ERA HOMOLOG	96	91	912
94	3	951	2741	gi 153615	phosphoenolpyruvate:sugar phosphotransferase system enzyme I [Streptococcus salivarius]	96	92	1791
127	1	1	168	gi 1581599	Initiation factor IF-1 [Lactococcus lactis]	96	89	168
128	14	10438	11154	gi 1276873	pepO [Streptococcus thermophilus]	96	93	717
181	4	1362	1598	gi 46606	laciD polypeptide [AA 1-326] [Staphylococcus aureus]	96	80	237
218	1	1	834	gi 1743856	Intragenic coaggregation-relevant adhesin [Streptococcus gordonii]	96	93	834
319	2	115	441	gi 208225	heat-shock protein 82/neomycin phosphotransferase fusion protein (hsp82-neo)	96	96	327
54	12	8622	10967	gnl PBD 00972	Pyruvate formate-lyase [Streptococcus mutans]	95	89	2346
181	2	606	1289	gi 1149396	laciD [Lactococcus lactis]	95	89	684
46	3	3410	3045	gi 1850606	[1]XW [Streptococcus mutans]	94	86	366
89	10	7972	7337	gi 703442	thymidine kinase [Streptococcus gordonii]	94	94	636
148	9	6431	7354	gi 1995767	UDP-glucose pyrophosphorylase [Streptococcus pyogenes]	94	95	924
160	7	4430	5848	gi 153573	H ⁺ ATPase [Enterococcus faecalis]	94	87	1419
2	3	4598	3513	gi 153763	plasmin receptor [Streptococcus pyogenes]	93	86	1086
12	8	7877	6204	gi 1103065	formyl-tetrahydrofolate synthetase [Streptococcus mutans]	93	84	1674

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
65	11 4734	5120	g91 40150	[L14 protein (AA 1-122) [Bacillus subtilis]		93	87	387
68	1 53	1297	g91 47341	[antitumor protein [Streptococcus pyogenes]		93	87	1245
80	1 3	299	g91 PID0166	[ribosomal protein S7 [Bacillus subtilis]		93	84	297
127	3 695	1093	g91 142462	[ribosomal protein S11 [Bacillus subtilis]		93	86	399
160	5 1924	3462	g91 1773264	[ATPase, alpha subunit [Streptococcus mutans]		93	85	1539
211	5 3757	3047	g91 53273	[aminopeptidase C [Streptococcus thermophilus]		93	82	711
262	1 16	564	g91 149394	[lacB [Lactococcus lactis]		93	90	549
366	1 197	3	g91 295259	[tryptophan synthase beta subunit [Synechocystis sp. 1]		93	91	195
25	3 1392	1976	g91 1574496	[hypothetical [Haemophilus influenzae]		92	80	585
36	21 2081	19927	g91 310632	[hydrophobic membrane protein [Streptococcus gordonii]		92	86	855
181	3 1265	1534	g91 149396	[lacD [Lactococcus lactis]		92	83	270
181	7 3652	4060	g91 149410	[enzyme III [Lactococcus lactis]		92	83	399
32	4 5631	3937	g91 PID0294090	[fibronectin-binding protein A [Streptococcus gordonii]		91	85	1695
46	2 3054	1462	g91 1850607	[signal recognition particle Ffh [Streptococcus mutans]		91	84	1593
65	10 4742	4726	g91 S1785 S178	[ribosomal protein S17 - Bacillus stearothermophilus		91	80	285
77	2 260	1900	g91 287871	[groEL gene product [Lactococcus lactis]		91	82	1641
84	1 2	2056	g91 8771784	[Clp-like ATP-dependent protease binding subunit [Bos taurus]		91	79	2055
99	8 10550	9272	g91 153740	[sucrose phosphorylase [Streptococcus mutans]		91	84	1479
99	9 11947	11072	g91 153739	[membrane protein [Streptococcus mutans]		91	78	876
127	5 2065	2469	g91 S07223 RBS	[ribosomal protein L17 - Bacillus stearothermophilus		91	78	405
132	6 9339	9390	g91 142065	[hsp60 (Bacillus stearothermophilus)]		91	89	150
137	8 4765	6153	g91 PID0100347	[Na ⁺ -ATPase beta subunit [Enterococcus hirae]		91	79	1389
151	7 11119	9734	g91 1815634	[glutamine synthetase type 1 [Streptococcus agalactiae]		91	82	1386
201	2 1798	278	g91 2208938	[dextran glucosidase DexS [Streptococcus suis]		91	79	1521
222	2 673	1839	g91 153741	[ATP-binding protein [Streptococcus mutans]		91	85	1167
293	5 4113	4400	g91 1196931	[unknown protein [Insertion sequence IS861]		91	71	288
32	7 6166	6570	g91 A3693 A369	[diacylglycerol kinase homolog - streptococcus mutans]		90	77	405

TABLE 2 *S. pneumoniae* - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
33	2	841	527	gi 1196921	unknown protein [Insertion sequence IS861]	90	70	315
48	127	20508	19757	gnl PID e274705	lactate oxidase [Streptococcus iniae]	90	80	1152
55	121	19777	18515	gnl PID e221213	ClpX protein [Bacillus subtilis]	90	75	1263
56	2	717	977	gi 1710133	flagellar filament cap [Borrelia burgdorferi]	90	50	261
65	1	1	606	gi 1165303	L3 [Bacillus subtilis]	90	75	606
114	1	2	988	gi 153562	aspartate beta-semialdehyde dehydrogenase (EC 1.2.1.11) [Streptococcus mutans]	90	80	987
120	1	1345	827	gi 407880	ORF1 [Streptococcus equisimilis]	90	75	519
159	12	7630	8298	gi 143012	GMP synthetase [Bacillus subtilis]	90	84	609
166	4	4076	3282	gi 1661179	high affinity branched chain amino acid transport protein [Streptococcus mutans]	90	78	795
183	1	28	1395	gi 308858	ATP pyruvate 2-O-phosphotransferase [Lactococcus lactis]	90	76	1368
191	3	2891	1662	gi 149521	tryptophan synthase beta subunit [Lactococcus lactis]	90	78	1230
198	2	1551	436	gi 2323342	(AF014460) CCPA [Streptococcus mutans]	90	80	747
305	1	37	783	gi 157551	asparagine synthetase A (asnA) [Haemophilus influenzae]	89	78	1059
8	3	2285	3343	gi 149334	putative [Lactococcus lactis]	90	76	1116
46	8	7577	7362	gi 145434 A454	ribosomal protein L19 - Bacillus stearothermophilus	89	76	216
49	9	8363	10342	gi 153792	recP peptide [Streptococcus pneumoniae]	89	83	1980
51	14	18410	19447	gi 108877	ATP:D-fructose 6-phosphate 1-phosphotransferase [Lactococcus lactis]	89	81	1038
57	11	9686	10669	gnl PID d100932	H2O-forming NADH Oxidase [Streptococcus mutans]	89	77	984
65	5	2418	2786	gi 1165107	S19 [Bacillus subtilis]	89	81	369
65	8	3806	4225	lsp P14577 Ru16-	50S RIBOSOMAL PROTEIN L16.	89	82	420
65	18	8219	8719	gi 143417	ribosomal protein S5 [Bacillus stearothermophilus]	89	76	501
73	9	6337	5315	gi 532204	prS [Listeria monocytogenes]	89	70	1023
76	3	3360	1465	gnl PID e200671	lepa gene product [Bacillus subtilis]	89	76	1836
99	10	112818	111919	gi 1537378	membrane protein [Streptococcus mutans]	89	73	900
120	2	3552	1300	gi 407881	stringent response-like protein [Streptococcus equisimilis]	89	79	2253
122	5	4512	2791	gnl PID e280490	unknown [Streptococcus pneumoniae]	89	81	1722

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
176	1	669	4	[gi 47394	[5'-oxoprolyl-peptidase [Streptococcus pyogenes]	89	78	666
177	6	3050	3934	[gi 912423	[putative [Lactococcus lactis]	89	71	885
181	8	4033	5751	[gi 149411	[enzyme III [Lactococcus lactis]	89	80	1719
211	4	3149	2793	[gi 335273	[aminopeptidase C [Streptococcus thermophilus]	89	70	408
361	1	431	838	[gi 1196922	[unknown protein [Insertion sequence IS861]	88	78	1305
34	17	11839	10535	[sp P10053 SYTH S	[HISTIDYL-TRNA SYNTHETASE (EC 6.1.1.21) [HISTIDINE-TRNA LIGASE] (M1SRS)]	88	78	978
38	3	1646	2623	[gi 2058544	[putative ABC transporter subunit ComY [Streptococcus gordonii]	88	78	357
54	1	3	227	[gi P010120	[YsgU [Bacillus subtilis]	88	66	225
57	2	611	1468	[gi P1013943	[putative reductase 1 [Saccharomyces cerevisiae]	88	75	858
65	13	5497	6069	[pir A29102 RB85	[ribosomal protein L5 - Bacillus stearothermophilus]	88	75	573
65	20	9030	9500	[gi 2078381	[ribosomal protein L15 [Staphylococcus aureus]	88	83	471
78	3	3636	1108	[gi P10100781	[lyzyl]-aminopeptidase [Lactococcus lactis]	88	80	2529
106	12	12965	12054	[gi 017421	[putative heat shock protein HtpX [Streptococcus gordonii]	88	72	912
107	1	219	962	[gi P0139862	[putative acylneuraminate lyase [Clostridium tertium]	88	75	744
111	8	14073	10420	[gi 402363]	[RNA Polymerase beta-subunit [Bacillus subtilis]	88	74	3654
126	9	11306	12062	[gi P0139868	[unknown [Bacillus subtilis]	88	61	270
140	17	19143	18874	[gi 1573659	[H. influenzae predicted coding region H10619 [Haemophilus influenzae]	88	88	1116
144	1	394	555	[gi P01274705	[lactate oxidase [Streptococcus faeciae]	88	75	162
148	4	2723	3493	[gi 1591672	[phosphate transport system ATP-binding protein [Methanococcus jannaschii]	88	68	771
160	8	5853	6278	[gi 1772267	[ATPase, epsilon subunit [Streptococcus mutans]	88	65	426
177	4	1770	2885	[gi 149426	[putative [Lactococcus lactis]	88	72	1116
200	5	2387	2998	[gi 1196922	[unknown protein [Insertion sequence IS861]	88	74	528
211	6	4140	3613	[gi 535273	[aminopeptidase C [Streptococcus thermophilus]	88	75	1359
291	6	2017	3375	[gi P0100571	[adenylosuccinate synthetase [Bacillus subtilis]	88	78	378
319	4	658	317	[gi 603578	[serine/threonine kinase [Phytophthora capsici]	88	88	342
40	5	4353	4514	[gi 153672	[lactose repressor [Streptococcus mutans]	87	56	162

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
49	10	10660	10929	[gi 1196921]	[unknown protein (Insertion sequence IS861)]	87	72	270
65	7	3140	3808	[gi 1165309]	[S3 (Bacillus subtilis)]	87	73	669
65	15	6823	7039	[gi 1044978]	[ribosomal protein S8 (Bacillus subtilis)]	87	73	417
75	8	5411	6625	[gi 1877422]	[galactokinase (Streptococcus mutans)]	87	78	1215
80	2	703	2805	[gi 101166]	[elongation factor G (Bacillus subtilis)]	87	76	2103
82	1	541	248	[gi 1196921]	[unknown protein (Insertion sequence IS861)]	87	69	294
140	123	125033	23897	[gi 1044979]	[phenylalanyl-tRNA synthetase beta subunit (Bacillus subtilis)]	87	74	1137
214	14	10441	8516	[gi 12281305]	[glucosidase inhibited division protein homolog Gida (Bacillus subtilis)]	87	75	1926
220	2	2712	874	[gi 1044978]	[product highly similar to elongation factor EF-G (Bacillus subtilis)]	87	73	1869
260	4	2066	2389	[gi 1196921]	[unknown protein (Insertion sequence IS861)]	87	72	294
323	1	27	650	[gi 1897795]	[30S ribosomal Protein (Pediococcus acidilactici)]	87	73	624
357	1	154	1570	[gi 1044978]	[ribosomal protein S8 (Bacillus subtilis)]	87	73	417
49	11	10927	11445	[gi 1196922]	[unknown protein (Insertion sequence IS861)]	86	63	519
59	12	7461	9224	[gi 1951051]	[relaxase (Streptococcus pneumoniae)]	86	68	1764
65	4	1553	2401	[gi 1027591RSBS]	[ribosomal protein L2 - Bacillus stearothermophilus]	86	77	849
65	123	10957	11610	[gi 44074]	[adenylate kinase (Lactococcus lactis)]	86	76	654
82	4	4374	4856	[gi 153745]	[mannitol-specific enzyme III (Streptococcus mutans)]	86	72	483
102	4	4270	4986	[gi 10432]	[OMP decaboxylase (Lactococcus lactis)]	86	76	717
106	6	7824	6880	[gi 10137598]	[aspartate transcarbamylase (Lactobacillus leichmannii)]	86	71	273
107	1	1	273	[gi 10339862]	[putative acylneuraminate lyase (Clostridium tertium)]	86	73	1531
111	7	10432	6710	[gi 10264705]	[OMP-dependent RNA polymerase (Streptococcus pyogenes)]	86	80	3723
131	9	5704	4892	[gi 1661193]	[polypeptide diacylglycerol transferase (Streptococcus mutans)]	86	71	813
134	7	6430	7980	[gi 2388637]	[glycerol kinase (Enterococcus faecalis)]	86	72	891
146	11	7473	6583	[gi 1591731]	[malvalonate kinase (Methanococcus jannaschii)]	86	78	1416
153	2	595	2010	[gi 2160707]	[dipeptidase (Lactococcus lactis)]	86	74	1434
154	1	2	1435	[gi 1857246]	[6-phosphogluconate dehydrogenase (Lactococcus lactis)]	86	74	1434

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig ID	ORF ID	start (nt)	stop (nt)	match accession	match gene name	sim		sim	
						%	length (nt)	%	length (nt)
161	5	505	6284	[gi 47529]	[Unknown [Streptococcus salivarius]	86	66	86	1260
184	1	2	1483	[gi 642667]	[NAD+-dependent glyceraldehyde-3-phosphate dehydrogenase (Streptococcus mutans)]	86	73	86	1482
210	8	3639	6571	[gi 153661]	[translational initiation factor IF2 (Enterococcus faecium)]	86	76	86	2913
250	1	2	187	[gi 1573551]	[asparagine synthetase A (asnA) [Haemophilus influenzae]]	86	68	86	186
36	4	2614	3909	[gi 2149909]	[cell division protein (Enterococcus faecalis)]	85	73	85	1266
38	4	2475	3587	[gi 2058545]	[putative ABC transporter subunit ComBA (Streptococcus gordoni)]	85	72	85	1113
38	5	3577	3915	[gi 2058546]	[ComYC (Streptococcus gordoni)]	85	80	85	339
57	5	12997	3789	[gi 1203136]	[YqfF (Bacillus subtilis)]	85	72	85	993
82	5	4915	6054	[gi 153746]	[mannitol-phosphate dehydrogenase (Streptococcus mutans)]	85	68	85	1140
83	15	14690	15793	[gi 143371]	[phosphoribosyl aminomimidazole synthetase (PUR-H) [Bacillus subtilis]]	85	69	85	1104
87	2	14117	2388	[gi 1184967]	[ScrR (Streptococcus mutans)]	85	69	85	972
108	3	2666	3154	[gi 153366]	[ORF (19k protein) [Enterococcus faecalis]]	85	67	85	489
127	2	312	1692	[gi 1044989]	[ribosomal protein S13 (Bacillus subtilis)]	85	72	85	381
128	3	1534	2409	[gi 1685110]	[tetrahydrofolate dehydrogenase/cyclohydrolase (Streptococcus thermophilus)]	85	71	85	876
137	1	2962	4767	[gi 100347]	[Na+-ATPase alpha subunit (Enterococcus hirae)]	85	74	85	1806
170	2	2622	709	[gi 102006]	[AB001498] FUNCTION UNKNOWN, SIMILAR PRODUCT IN E.COLI, H. INFLUENZAE AND NEISSERIA HEMINGVITDIS. [Bacillus subtilis]	85	70	85	1914
187	5	3760	4386	[gi 1727436]	[putative 20-kDa protein [Lactococcus lactis]]	85	65	85	627
233	2	728	1873	[gi 1163116]	[ORF-5 (Streptococcus pneumoniae)]	85	67	85	1146
234	3	962	1255	[gi 2293155]	[IAP002220] YtaA (Bacillus subtilis)]	85	61	85	294
240	1	309	1931	[gi 143597]	[CTP synthetase [Bacillus subtilis]]	85	70	85	1623
6	1	199	1521	[gi 508979]	[GTP-binding protein [Bacillus subtilis]]	84	72	84	1323
19	4	1793	2593	[gi 2352484]	[RNaseH II (Lactococcus lactis)]	84	68	84	801
10	4	4375	3443	[gi 100584]	[putative acyli neuraminate lyase (Clostridium tertium)]	84	70	84	913
20	17	17720	19687	[gi 120584]	[cell division protein (Bacillus subtilis)]	84	71	84	1938
22	128	21723	20884	[gi 299163]	[alanine dehydrogenase (Bacillus subtilis)]	84	68	84	840

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
30	10	7730	6792	gnl PID d100296	fructokinase [Streptococcus mutans]	84	75	939
33	9	5650	5300	gi 147194	lpha protein [Escherichia coli]	84	71	351
36	122	121551	120772	gi 310631	ATP binding protein [Streptococcus gordonii]	84	72	780
48	4	2837	2505	gi 888269	6-phospho-beta-glucosidase [Escherichia coli]	84	69	333
58	1	41	1516	gi 450849	anylyase [Streptococcus bovis]	84	73	1476
59	10	6715	7116	gi 951033	[ORF10] putative [Streptococcus pneumoniae]	84	74	402
62	3	21	644	gi 806487	[ORF211] putative [Lactococcus lactis]	84	66	624
65	17	7779	8207	gi 1044980	ribosomal protein L18 [Bacillus subtilis]	84	73	429
65	21	9507	10397	gi 144073	ISCEY protein [Lactococcus lactis]	84	68	891
106	4	5474	2262	gnl PID e159387	[carbamoyl]-phosphate synthase [Lactobacillus plantarum]	84	73	3213
159	1	147	4	gi 806487	[ORF211] putative [Lactococcus lactis]	84	63	144
163	4	4690	5910	gi 12293164	[AF008220] SAM synthase [Bacillus subtilis]	84	69	1221
192	1	46	1308	gi 495046	tripeptidase [Lactococcus lactis]	84	73	1263
348	1	671	6	gi 1787753	[AE000245] f346; 79 pct identical to 336 amino acids of Adh1_ZYMO SW: P20368 but has 10 additional N-ter residues [Escherichia coli]	84	71	666
3	4	1572	3575	gi 143766	[Itahsv] (EC 6.1.1.3) [Bacillus subtilis]	83	65	2004
9	6	3893	3417	gnl PID d100576	single strand DNA binding protein [Bacillus subtilis]	83	68	477
17	15	7426	8457	gi 520738	[comA] protein [Streptococcus pneumoniae]	83	66	1032
20	12	13860	14144	gnl PID d100583	lumkown [Bacillus subtilis]	83	61	285
23	4	3358	2606	gi 1788294	[AE000290] o238; This 238 aa orf is 40 pct identical (5 gaps) to 231 residues of an approx. 248 aa protein YEC_ECOLI SW: P24337 [Escherichia coli]	83	74	753
28	6	3304	3005	gi 1571659	H. influenzae predicted coding region H10559 [Haemophilus influenzae]	83	57	300
35	7	5108	3867	gi 1311707	hypothetical nucleotide binding protein [Archoplasmalaidlawii]	83	63	1242
55	19	17932	17528	gi 537085	[ORF_f14] [Escherichia coli]	83	59	405
55	20	18539	17919	gi 496258	[orfX] [Bacillus subtilis]	83	69	621
65	6	2795	3142	gi 1165308	[L22] [Bacillus subtilis]	83	64	348
68	6	6877	6683	gi 1212194	immunoglobulin A1 protease [Streptococcus pneumoniae]	83	54	195

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF	Start	Stop	match	match gene name	% sim	% ident	length
ID	ID	(nt)	(nt)	accession				(nt)
87	15	15112	14771	[gnl PID d123522]	[putative rpoZ protein (Bacillus subtilis)]	83	54	312
96	12	8963	9631	[gi 47394]	[5'-oxoprolyl-peptidase (Streptococcus pyogenes)]	83	73	659
98	1	3	263	[gi 1183865]	[glutamine-binding subunit (Bacillus subtilis)]	83	55	261
120	4	7170	5233	[gi 310630]	[zinc metalloprotease (Streptococcus gordonii)]	83	72	1938
127	7	2998	4347	[gi 1500567]	[M. jannaschii predicted coding region #A1665 (Methanococcus jannaschii)]	83	72	1350
137	1	3	440	[gi 472918]	[v-type Na-ATPase (Enterococcus hirae)]	83	60	438
160	6	3466	4356	[gi 1773265]	[ATPase, gamma subunit (Streptococcus mutans)]	83	67	891
214	4	2278	2964	[gi 663229]	[transposase (Streptococcus pneumoniae)]	83	72	687
226	3	2367	2020	[gi 142154]	[thioredoxin (Synchococcus PCC6301)]	83	58	148
303	1	3	1049	[gi 40046]	[phosphoglucose isomerase A (AA 1-449) (Bacillus stearothermophilus)]	83	67	1047
303	2	1155	1931	[gi 289282]	[glutamyl-tRNA synthetase (Bacillus subtilis)]	83	67	777
6	17	15370	14318	[gi 333147]	[ribose-phosphate pyrophosphokinase (Bacillus caldolyticus)]	82	64	1053
7	1	299	96	[gi 143668]	[ribosomal protein L28 (Bacillus subtilis)]	82	69	204
9	3	1479	1090	[gi 1885178]	[unknown (Bacillus subtilis)]	82	46	390
9	7	4213	3899	[gnl PID d100576]	[ribosomal protein S6 (Bacillus subtilis)]	82	60	315
12	6	4688	3942	[gnl PID d100571]	[unknown (Bacillus subtilis)]	82	68	747
22	17	13422	14837	[gi 520754]	[putative (Bacillus subtilis)]	82	69	1416
22	18	14897	15658	[gnl PID d101929]	[uridine monophosphate kinase (Synochocystis sp.)]	82	62	762
33	16	11471	10641	[gnl PID d101190]	[ORF4 (Streptococcus mutans)]	82	68	831
35	9	7400	6255	[gi 1881543]	[UDP-N-acetylglucosamine-2-epimerase (Streptococcus pneumoniae)]	82	68	1146
40	10	8003	7533	[gi 1173519]	[riboflavin synthase beta subunit (Actinobacillus pleuropneumoniae)]	82	68	471
48	32	23159	23437	[gi 1930092]	[outer membrane protein (Campylobacter jejuni)]	82	61	279
52	14	13833	14755	[gi 142321]	[deoxyribodipyridine photolyase (Bacillus subtilis)]	82	61	933
60	4	4737	1849	[gnl PID d02221]	[Dalnococcus radiodurans]	82	66	2889
62	4	2131	1457	[gi 2246749]	[AF009622) thioredoxin reductase (Listeria monocytogenes)]	82	63	675
71	11	16586	17518	[gnl PID e322053]	[ss-1,4'-galactosyltransferase (Streptococcus pneumoniae)]	82	60	913
73	13	9222	7837	[gnl PID d100566]	[unknown (Bacillus subtilis)]	82	65	1386

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
74	1	1	3771	[gnl PTD 010199]	alkaline amylopullulanase [Bacillus sp.]	82	68	3771
83	9	1696	3983	[gnl PTD 0305362]	unnamed protein product [Streptococcus thermophilus]	82	52	288
86	11	10776	9394	[gi 6833583]	[5'-enolpyruvyl]shikimate-3'-phosphate synthase [Lactococcus lactis]	82	67	1383
89	12	8295	9752	[gi 140025]	homologous to E. coli 50K [Bacillus subtilis]	82	66	1458
115	9	10347	8812	[gnl PTD 02090]	[AB003927] phospho-beta-galactosidase 1 [Lactobacillus gasseri]	82	74	1536
118	1	1	1332	[gnl PTD 00579]	seryl-tRNA synthetase [Bacillus subtilis]	82	71	1332
151	3	4657	6246	[pir S0697 S060]	type I site-specific deoxyribonuclease (EC 3.1.21.3) CtrA chain S - Citrobacter freundii	82	66	1550
173	6	4183	3503	[gi 2313836]	[AE000584] conserved hypothetical protein [Helicobacter pylori]	82	68	681
177	12	5481	7442	[gnl PTD 0101999]	[AB01341] NcrB [Escherichia coli]	82	58	1962
193	2	178	576	[pir S08564 RBS S08564]	ribosomal protein S9 - Bacillus stearothermophilus	82	70	399
245	2	258	845	[gi 146402]	[EC04] type I restriction-modification enzyme S subunit [Escherichia coli]	82	68	588
9	5	3400	3146	[gnl PTD 00576]	ribosomal protein S18 [Bacillus subtilis]	81	66	255
16	7	7484	8413	[gi 1100074]	[tryptophanyl]-tRNA synthetase [Clostridium longisporum]	81	63	3513
20	11	10308	13820	[gnl PTD 00583]	transcription-repair coupling factor [Bacillus subtilis]	81	63	375
38	2	1232	1606	[gi 12058543]	[putative DNA binding protein [Streptococcus gordoni]]	81	67	1311
45	2	3061	1751	[gi 460259]	lenoase [Bacillus subtilis]	81	61	1266
46	1	2	1267	[gi 431231]	luracil permease [Bacillus caldolyticus]	81	70	1014
48	3	2453	1440	[gnl PTD 00453]	[mannosephosphate isomerase [Streptococcus mutans]]	81	64	771
54	2	1106	336	[gi 154752]	transport protein [Agrobacterium tumefaciens]	81	66	516
65	122	10306	10821	[gi 44073]	SecY protein [Lactococcus lactis]	81	69	1272
89	4	3874	2603	[gi 556886]	serine hydroxymethyltransferase [Bacillus subtilis]	81	75	198
99	16	19126	18929	[gi 2313526]	[AE000557] H. pylori predicted coding region HP0411 [Halicobacter pylori]	81	61	552
106	7	8373	7822	[gnl PTD 039384]	[pyrR [Lactobacillus plantarium]]	81	66	1824
108	6	5054	6877	[gi 146939]	group B oligopeptidase PepB [Streptococcus agalactiae]	81	65	2305
113	15	15899	18283	[pir S09411 S094]	sporII protein - Bacillus subtilis	81	69	276
128	5	3359	3634	[gi 1685111]	[lort109] [Streptococcus thermophilus]	81	+	+

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
10	1	830	3211	[gi 304896	Eco type I restriction-modification enzyme R subunit [Escherichia coli]	81	59	2382
151	1	6722	7837	[gi 1223288	GMP synthetase [Bacillus subtilis]	81	69	1116
159	1	739	458	[gi 19001488]	FUNCTION UNKNOWN. [Bacillus subtilis]	81	55	282
170	1	1759	893	[gi 149522	tryptophan synthase alpha subunit [Lactococcus lactis]	81	65	867
191	2	1	1994	[gi 1575787	reverse transcriptase endonuclease [Drosophila virilis]	81	43	297
214	3	2290	4	[gi 1415	cellulose phosphotransferase enzyme II. [Bacillus stearothermophilus]	81	59	408
217	4	4008	4008	[gi 146673	tryptophan synthase beta subunit [Lactococcus lactis]	81	68	300
262	2	569	868	[gi 153675	tagatose 6-P kinase [Streptococcus mutans]	81	60	660
299	1	1663	4	[gi 1900154	StySRI methylase [Salmonella enterica]	81	59	294
366	2	376	93	[gi 149521	tryptophan synthase beta subunit [Lactococcus lactis]	80	64	477
12	110	8766	9242	[gi 11216490	DNA/pantothenate metabolism flavoprotein [Streptococcus mutans]	80	64	477
17	111	6050	5748	[gi 19001562]	unnamed protein product [Streptococcus thermophilus]	80	67	303
17	116	8455	9066	[gi 1703126	leucocin A translocator [Leuconostoc gelidum]	80	59	612
18	13	2440	1613	[gi 1591672	phosphate transport system ATP-binding protein [Methanococcus jannaschii]	80	58	828
27	3	4248	1579	[gi 1452209	valyl-tRNA synthetase [Bacillus subtilis]	80	69	2670
28	7	3671	3288	[gi 1573760	H. influenzae predicted coding region H10660 [Haemophilus influenzae]	80	63	384
32	2	902	1933	[gi 19001549]	dihydroorotate dehydrogenase B [Lactococcus lactis]	80	66	1032
39	1	1	1266	[gi 19001578]	hom [Lactococcus lactis]	80	63	1266
52	5	4363	3593	[gi 1183884	ATP-binding subunit [Bacillus subtilis]	80	57	771
54	5	4550	4744	[gi 2198820	Cux/CDP (1B1); Cux/CDP homoprotein [Mus musculus]	80	60	195
59	11	7109	7486	[gi 1951052	[AF004225] ORF9, putative [Streptococcus pneumoniae]	80	68	378
65	3	1230	1550	[pir A02815]R5BS	ribosomal protein L23 - Bacillus stearothermophilus	80	69	321
65	12	5174	5503	[pir A02819]R5BS	ribosomal protein L24 - Bacillus stearothermophilus	80	70	330
66	9	9884	10687	[gi 2313836	[AE000584] conserved hypothetical protein [Helicobacter pylori]	80	66	804
82	2	648	2438	[gi 1622991	mannitol transport protein [Bacillus stearothermophilus]	80	65	1791
85	1	950	630	[gi 1528955	polyketide synthase [Bacillus subtilis]	80	46	321
89	8	6870	5779	[gi 853776	peptide chain release factor 1 [Bacillus subtilis]	80	63	1092
93	12	8718	7438	[gi 19001599]	hypothetical protein [Synchrocoystis sp.]	80	60	1281

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
106	5	6854	5751	gnl PID e199386	glutaminase of carbamoyl-phosphate synthase [Lactobacillus plantarum]	80	65	1104
109	2	2160	1450	gi 40056	phoP gene product [Bacillus subtilis]	80	59	711
124	9	4246	3953	gnl PID d102254	[30S ribosomal protein S16 [Bacillus subtilis]	80	65	294
128	8	5148	6428	gi 2281308	phosphopentomutase [Lactococcus lactis cremoris]	80	66	1281
137	19	16665	11376	gi 159109	[NADP-dependent glutamate dehydrogenase [Giardia intestinalis]	80	68	1290
140	19	15699	19457	gi 517210	porative transposase [Streptococcus pyogenes]	80	70	243
158	2	2474	984	gi 1877423	galactose-1-p-uridyl transferase [Streptococcus mutans]	80	65	1491
171	10	7474	7728	gi 397800	cyclophilin C-associated protein [Mus musculus]	80	60	255
181	1	2	619	gi 149395	lacc [Lactococcus lactis]	80	66	618
313	1	27	539	gi 143467	[ribosomal protein S4 [Bacillus subtilis]	80	70	513
329	2	1652	858	gi 533080	ReEF protein [Streptococcus pyogenes]	80	63	795
371	1	2	958	gi 442360	CIPC adenosine triphosphatase [Bacillus subtilis]	80	58	957
8	7	4312	5580	gi 149435	[putative [Lactococcus lactis]	79	64	1269
23	1	1175	135	gi 1542975	AbcB [Thermoaerobacterium thermosulfurigenes]	79	61	1041
33	14	9244	8201	gnl PID e253891	[UDP-glucose 4 epimerase [Bacillus subtilis]	79	62	1044
36	3	1242	2633	gnl PID e324218	[ftcA [Enterococcus hirae]	79	58	1392
38	13	7155	8378	gi 405134	acetate kinase [Bacillus subtilis]	79	58	1224
55	7	9011	8229	gi 1146234	[dihydrodipicolinate reductase [Bacillus subtilis]	79	56	783
65	19	8661	8915	gi 2078380	[ribosomal protein L10 [Staphylococcus aureus]	79	68	255
69	4	3678	2128	gnl PID e311452	[unknown [Bacillus subtilis]	79	64	1551
69	9	7881	7279	gi 1677850	[hypothetical protein [Staphylococcus aurous]	79	59	603
72	10	8491	9783	gnl PID d101091	[hypothetical protein [Synchocystis sp.]	79	62	1293
80	3	2906	7300	gi 143342	[polymerase III [Bacillus subtilis]	79	65	4395
82	14	133226	15689	gnl PID e255093	[hypothetical protein [Bacillus subtilis]	79	65	2164
86	13	112233	11118	gi 683582	[prephenate dehydrogenase [Lactococcus lactis]	79	58	1116
92	3	940	1734	gi 537286	[triosephosphate isomerase [Lactococcus lactis]	79	65	795
98	6	4023	4742	gnl PID d100262	[Lys protein [Salmonella typhimurium]	79	63	720

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
99	12	16315	14150	gi 1153736	α -galactosidase [Streptococcus mutans]	79	64	2166
107	7	5884	6406	gi 460080	D-alanine:D-alanine ligase-related protein [Enterococcus faecalis]	79	58	723
113	9	6958	8303	gi 466882	[ppb1; B1496_C2_189] [Mycobacterium leprae]	79	64	1446
151	10	1324	12213	gi 450686	[3'-phosphoglycerate kinase [Thermotoga maritima]	79	60	1212
162	2	1158	3017	gi 506700	[Capp1; Staphylococcus aureus]	79	67	1860
177	5	2876	3052	gi 912423	[putative [Lactococcus lactis]	79	61	177
177	8	4198	4563	gi 1169429	[putative [Lactococcus lactis]	79	61	366
187	3	2228	2907	gi 1402002	[AB001488] FUNCTION UNKNOWN [Bacillus subtilis]	79	53	180
189	7	3589	4350	gi 143449	putative ATP-binding protein of ABC-type [Bacillus subtilis]	79	61	762
191	5	4249	3449	gi 1149519	[indoleglycerol phosphate synthase [Lactococcus lactis]	79	66	801
211	3	1805	2737	gi 147404	[mannose permease subunit II-M-Man [Escherichia coli]	79	57	933
212	1	1863	3621	gi 140209004	[glutaredoxin-like protein [Lactococcus lactis]	79	58	243
215	1	987	715	gi 1293212	[ATP08220] arginine succinate synthase [Bacillus subtilis]	79	64	273
323	2	530	781	gi 837795	[30S ribosomal protein [Pediococcus acidilactici]	79	67	252
380	1	694	2	gi 118460	[polynucleotide phosphorylase [Bacillus subtilis]	79	64	693
384	2	655	239	gi 143328	[phop protein (put.); putative [Bacillus subtilis]	79	59	417
6	3	2820	4091	gi 853767	[UDP-N-acetylglucosamine 1-carboxyvinyltransferase [Bacillus subtilis]	78	62	1272
8	1	50	1786	gi 149432	[putative [Lactococcus lactis]	78	63	1737
9	1	351	124	gi 897793	[y98 gene product [Pediococcus acidilactici]	78	59	228
15	8	7364	8314	gi 1400585	[cysteine synthetase A [Bacillus subtilis]	78	63	951
20	10	9738	10310	gi 1400583	[stage V sporulation [Bacillus subtilis]	78	58	573
20	16	117165	117113	gi 49105	[hypoxanthine phosphoribosyltransferase [Lactococcus lactis]	78	59	549
22	22	117388	118416	gi 1401315	[YQE1; Bacillus subtilis]	78	60	1029
22	27	120371	120612	gi 1299163	[alanine dehydrogenase [Bacillus subtilis]	78	59	360
34	8	7407	7105	gi 41015	[aspartate-tRNA ligase [Escherichia coli]	78	55	303
35	8	6557	5196	gi 1657614	[Cap8E [Staphylococcus aureus]	78	60	1052

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start ID	Stop ID	match accession	match gene name	% sim	% ident	length (nt)
40	11	9287	8001	gi 1173518	GTP cyclohydrolase II/ 3,4-dihydroxy-2-butanone-4-phosphate synthase [Actinobacillus Pleuropneumoniae]	78	58	1287
48	31	22422	23183	gi 2314330	(AE000623) glutamine ABC transporter, ATP-binding protein [glnQ] [Helicobacter pylori]	78	58	762
52	2	2101	1430	gi 1163887	integral membrane protein [Bacillus subtilis]	78	54	672
55	14	13605	12712	gi 117326	(AB002150) YbbP [Bacillus subtilis]	78	58	894
55	17	16637	15612	gi 1173027	hypothetical protein [Bacillus subtilis]	78	51	1026
71	14	19756	19598	gi 1179164	calcium channel alpha-1D subunit [Homo sapiens]	78	57	159
74	111	15031	14018	gi 11573279	Holliday junction DNA helicase [UvB] [Haemophilus influenzae]	78	57	1014
75	9	6623	7972	gi 11877423	galactose-1-P-uridylyl transferase [Streptococcus mutans]	78	62	1350
81	12	12125	113906	gi 11573607	D-fucose isomerase (fucI) [Haemophilus influenzae]	78	66	1782
82	3	2423	4417	gi 1153744	ORF X: putative [Streptococcus mutans]	78	64	1995
83	18	16926	18500	gi 1143373	phosphoribosyl aminonimidazole carboxylase formyl formyltransferase/inosine monophosphate cyclohydrolase (PUR-HJ) [Bacillus subtilis]	78	63	1575
83	20	120212	120775	gi 1143364	phosphoribosyl aminonimidazole carboxylase I (PUR-E) [Bacillus subtilis]	78	64	564
92	2	165	878	gi 1173188	ORF2 [Streptococcus mutans]	78	62	714
98	8	5863	6909	gi 11731287	(AF013188) release factor 2 [Bacillus subtilis]	78	63	1047
113	3	1071	2741	gi 1580914	dnazX [Bacillus subtilis]	78	64	1671
127	4	1133	2071	gi 1142463	RNA polymerase alpha-core-subunit [Bacillus subtilis]	78	59	939
132	1	2782	497	gi 1561763	pullulanase [Bacteroides thetaiotaomicron]	78	58	2286
135	4	2698	3537	gi 1788036	(AE000269) NH3-dependent NAD synthetase [Escherichia coli]	78	66	840
140	124	126853	125423	gi 1100077	phospho-beta-glucosidase [Clostridium longi] sporum	78	64	1431
150	5	4690	4514	gi 149464	amino peptidase [Lactococcus lactis]	78	42	177
152	1	1	795	gi 639915	NADH dehydrogenase subunit [Thunbergia alata]	78	43	795
162	4	4997	4110	gi 11733528	putative YbaP protein [Bacillus subtilis]	78	64	888
181	10	8651	7947	gi 149402	lactose repressor (lacR; alt.) [Lactococcus lactis]	78	48	705
200	4	3627	4958	gi 11730172	Invertase [Zymomonas mobilis]	78	61	1332
203	3	3230	3015	gi 1174237	CycK [Pseudomonas fluorescens]	78	57	216

TABLE 2 *S. pneumoniae* - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	start (nt)	stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
210	9	6789	7172	gi 580902	[Bacillus subtilis] ORF6 gene product [Bacillus subtilis]	78	42	384
214	6	1810	2797	gnl PID0102049	[P. haemolytica] o-sialoglycoprotein endopeptidase; P36175 (660)	78	60	1014
214	13	6322	8163	gi 137781	[Bacillus subtilis] unknown [Bacillus subtilis]	78	62	1842
217	1	9	2717	gi 48830	[Intamoeba histolytica] alcohol dehydrogenase 2	78	64	2709
222	3	2316	3098	gi 1573047	[spore germination and vegetative growth protein (gerC2) [Haemophilus influenzae]]	78	65	783
268	1	742	8	gi 517210	[putative transposase [Streptococcus pyogenes]]	78	65	735
276	1	223	753	gnl PID00306	[ribosomal protein L1 [Bacillus subtilis]]	78	65	531
312	3	1567	1079	gi 1289261	[comE ORF2 [Bacillus subtilis]]	78	54	489
339	1	117	794	gi 1916729	[cadD [Staphylococcus aureus]]	78	53	678
342	2	762	265	gi 1842439	[phosphatidylglycerophosphate synthase [Bacillus subtilis]]	78	59	498
383	1	737	3	gi 1184680	[polynucleotide phosphorylase [Bacillus subtilis]]	78	64	735
7	15	11923	11018	gi 1398855	[carboxytransferase beta subunit [Synechococcus PCC7942]]	77	63	906
8	2	1698	2255	gi 149433	[putative [Lactococcus lactis]]	77	59	558
17	14	6948	7550	gi 520738	[comA protein [Streptococcus pneumoniae]]	77	60	603
30	12	9761	8967	gi 1000451	[Trep [Bacillus subtilis]]	77	43	795
36	14	11421	112131	gi 1573766	[phosphoglycromatase (gpmA) [Haemophilus influenzae]]	77	64	711
55	3	3836	4096	gi 1708640	[YeaB [Bacillus subtilis]]	77	55	261
61	8	8377	8054	gi 1890649	[multidrug resistance protein LmrA [Lactococcus lactis]]	77	51	324
65	2	607	1254	gi 40103	[ribosomal protein L4 [Bacillus stearothermophilus]]	77	63	648
68	8	7509	7240	gi 47551	[Mmp [Streptococcus suis]]	77	68	270
69	1	1083	118	gnl PID01311493	[unknown [Bacillus subtilis]]	77	57	966
77	5	4583	4026	gnl PID01281578	[hypothetical 12.2 kd protein [Bacillus subtilis]]	77	60	598
83	14	13104	14552	gi 1590947	[amidophosphotidylyltransferase [Methanococcus jannaschii]]	77	56	1449
94	4	3006	5444	gnl PID0132895	[AJ0004961] cyclic nucleotide-gated channel beta subunit [Rattus norvegicus]	77	66	2439
96	11	8518	8880	gi 251879	[ORF 1 [Lactococcus lactis]]	77	62	363
99	111	14082	12799	gi 153737	[sugar-binding protein [Streptococcus mutans]]	77	61	1284

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
105	2	361	1176	[gi]118921	[LysC protein (Haemophilus influenzae)]	77	51	816
108	4	3152	4030	[gi]1574730	[tetrurite resistance protein (tehB) (Haemophilus influenzae)]	77	58	879
110	4	3520	3131	[gi]1573900	[D-alanine permease (dagB) (Haemophilus influenzae)]	77	57	390
124	4	1796	1071	[gi]1573162	[tRNA (guanine-N1)-methyltransferase (trmD) (Haemophilus influenzae)]	77	58	726
126	4	5909	4614	[gnl]PID10163	[s2b (Bacillus subtilis)]	77	62	1296
128	2	630	1373	[gnl]PID101328	[ygi2 (Bacillus subtilis)]	77	58	744
130	1	1	1287	[gnl]PID102013	[hypothetical protein (Bacillus subtilis)]	77	61	1287
139	5	4388	3639	[gi]2291302	[AF0082201 YQQA (Bacillus subtilis)]	77	59	750
140	111	109311	9582	[gi]289284	[cysteinyl-tRNA synthetase (Bacillus subtilis)]	77	64	1350
140	18	18451	19263	[gi]517210	[putative transposase (Streptococcus pyogenes)]	77	66	189
141	2	976	1683	[gnl]PID10157887	[unf5 (aa 1-53) (Drosophila yakuba)]	77	50	708
141	4	2735	5293	[gi]556258	[secA (Listeria monocytogenes)]	77	59	2559
144	2	671	2173	[gnl]PID100585	[lysyl-tRNA thynthetase (Bacillus subtilis)]	77	61	1503
163	5	6412	7398	[gi]511015	[dihydroorotate dehydrogenase A (Lactococcus lactis)]	77	62	987
164	10	7841	7074	[gnl]PID100964	[homologue of iron dicitrate transport ATP-binding protein FecE of E. coli]	77	52	768
191	8	7257	5791	[gi]149516	[anthranilate synthase alpha subunit (Lactococcus lactis)]	77	57	1467
198	8	5377	5177	[gi]1573056	[hypothetical (Haemophilus influenzae)]	77	66	201
213	1	202	462	[gi]1743160	[BcrA2 (Mus musculus)]	77	50	261
250	2	231	509	[gnl]PID10334776	[YrbH protein (Bacillus subtilis)]	77	60	279
289	3	1737	1276	[gnl]PID100947	[Ribosomal Protein L10 (Bacillus subtilis)]	77	62	462
292	2	1399	668	[gi]143004	[transfer RNA-Gln synthetase (Bacillus stearothermophilus)]	77	58	732
7	3	2734	1166	[gnl]PID101824	[peptide-chain-release factor 3 (Synechocystis sp.)]	76	53	1559
7	23	18474	18235	[gi]455157	[acyl carrier protein (Cryptomonas ph1)]	76	57	240
9	8	5706	4342	[gi]1146247	[asparaginyl-tRNA synthetase (Bacillus subtilis)]	76	61	1365
10	5	4531	4185	[gnl]PID1034495	[hypothetical protein (Clostridium perfringens)]	76	53	147
18	2	1615	842	[gi]1591672	[phosphate transport system ATP-binding protein (Methanococcus jannaschii)]	76	56	774

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
22	37	27796	28173	[gnl PID e13389	translation initiation factor IF3 (AA 1-172) [Bacillus stearothermophilus]	76	64	378
35	6	3869	2682	[gi 1773346	[CaspG (Staphylococcus aureus)]	76	61	1188
48	28	21113	21787	[gi 2314328	[AE000623] glutamine ABC transporter, permease protein (glnP) [Helicobacter pylori]	76	52	675
52	12	12681	13786	[gi 142521	[deoxyribodipyrimidine photolyase [Bacillus subtilis]	76	58	906
55	10	11921	10571	[gnl PID e283110	fanD [Staphylococcus aureus]	76	61	951
57	8	7624	6559	[gi 1290561	[f0188 (Escherichia coli)]	76	47	1266
62	5	2406	2095	[gnl PID e131024	hypothetical protein [Bacillus subtilis]	76	59	312
65	9	4223	4441	[gi 40148	[L29 protein (AA 1-66) [Bacillus subtilis]	76	58	219
68	2	1328	12371	[gnl PID e284233	anabolic ornithine carboxyltransferase [Lactobacillus plantarum]	76	61	1044
69	8	7297	6005	[gnl PID d101420	[Pyrimidine nucleoside phosphorylase [Bacillus stearothermophilus]	76	61	1293
73	12	76319	7267	[gnl PID e243629	[unknown [Mycobacterium tuberculosis]]	76	53	573
74	5	8433	7039	[gnl PID d102048	[C. thermocellum beta-glucosidase: P16208 (1985) [Bacillus subtilis]]	76	60	1395
80	5	7643	7936	[gi 12314030	[ASD00599] conserved hypothetical protein [Helicobacter pylori]	76	61	294
82	15	16019	16596	[gi 1573900	[D-alanine permease (dagA) [Haemophilus influenzae]	76	56	978
83	19	18616	19884	[gi 143374	phosphoribosyl glycaminamide synthetase (PUR-D; gnt start codon) [Bacillus subtilis]	76	60	1269
87	16	15754	15110	[gnl PID e123500	putative Omk protein [Bacillus subtilis]	76	58	1179
86	14	13409	12231	[gi 143806	[AroF [Bacillus subtilis]	76	59	1440
87	1	1	3	[gi 1573804	[sucrose-6-phosphate hydrolase [Streptococcus mutans]	76	56	645
93	4	1769	1539	[gi 1574820	[1,4-alpha-glucan branching enzyme (g1gb) [Haemophilus influenzae]]	76	46	231
94	1	51	365	[gi 144313	[6.0 kd ORF [Plasmid ColE1]]	76	73	315
116	2	2151	1678	[gi 153641	[pneumococcal surface protein A [Streptococcus pneumoniae]]	76	59	474
123	6	3442	5895	[gi 1314297	[CtpC ATPase [Listeria monocytogenes]]	76	59	2454
126	2	2156	2932	[gnl PID d101328	[Yqz [Bacillus subtilis]]	76	61	77
128	10	6973	7797	[gi 1944944	[purine nucleoside phosphorylase [Bacillus subtilis]]	76	60	825
131	11	6186	5812	[gi 1674110	[AE000038] Mycoplasma pneumoniae, M2085 homolog, from M. genitalium [Mycoplasma pneumoniae]	76	47	375

TABLE 2 *S. pneumoniae* - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim			length (nt)
						% ident	% sim	% ident	
139	4	3641	3192	gi 12293302	[AF008220] Ytqa [Bacillus subtilis]	76	53	450	
140	114	14872	12536	gi 1184680	[polynucleotide phosphorylase [Bacillus subtilis]	76	62	2337	
143	2	2583	3905	gi 143795	[transfer RNA-Tyr synthetase [Bacillus subtilis]	76	61	1323	
170	6	5095	6114	gnl PID1d100959	[ycgQ [Bacillus subtilis]	76	44	1020	
180	2	1927	557	gi 40059	[orf 821 (aa 1-821) [Bacillus subtilis]	76	53	1371	
191	7	5815	5228	gi 551880	[lenthranilate synthase beta subunit [Lactococcus lactis]	76	61	588	
195	3	3829	2444	gi 2149905	[D-glutamic acid adding enzyme [Enterococcus faecalis]	76	60	1386	
200	3	1914	1629	gi 431272	[lysis protein [Bacillus subtilis]	76	58	1716	
201	1	431	207	gi 220898	[dextran glucosidase DexS [Streptococcus suis]	76	57	225	
214	2	1283	2380	gi 163328	[transposase [Streptococcus pneumoniae]	76	55	1098	
225	3	2338	3411	gi 1552775	[ATP-binding protein [Escherichia coli]	76	56	1074	
233	1	2	724	gi 1163115	[neuramidinase B [Streptococcus pneumoniae]	76	60	723	
347	1	523	38	gi 1537033	[orf_F156 [Escherichia coli]	76	60	486	
356	2	842	165	gi 2149905	[D-glutamic acid adding enzyme [Enterococcus faecalis]	76	61	678	
366	3	734	348	gi 149520	[phosphoribosyl anthranilate isomerase [Lactococcus lactis]	76	69	387	
5	8	112599	11484	gi 157493	[fimbrial transcription regulation repressor (pilB) [Haemophilus influenzae]	75	61	1116	
6	13	12553	11894	gnl PID1d102050	[ydhH [Bacillus subtilis]	75	51	660	
9	110	7282	6062	gi 142538	[aspartate aminotransferase [Bacillus sp.]	75	55	1221	
10	12	9080	7940	gi 149493	[scrFI methylase [Lactococcus lactis]	75	56	141	
18	5	4266	3301	gnl PID1d101319	[Yqgh [Bacillus subtilis]	75	52	966	
22	4	1838	2728	gi 1373157	orf-X; hypothetical protein; Method: conceptual translation supplied by author [Bacillus subtilis]	75	62	891	
30	11	9015	7828	gi 153801	[enzyme scr-II [Streptococcus mutans]	75	64	1188	
31	5	2362	2030	gi 2293211	[AF008220] putative thioredoxin [Bacillus subtilis]	75	53	333	
32	9	7484	8359	gnl PID1d100560	[formamidopyrimidine-DNA glycosylase [Streptococcus mutans]	75	61	876	
33	4	1735	1448	gi 4133976	[ipa-S2r gene product [Bacillus subtilis]	75	53	288	
33	10	670	5769	gi 5233105	[unknown [Bacillus subtilis]	75	56	702	

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
33	12	6678	7183	pir 00205 FECI	ferredoxin [Fe-4S] - <i>Clostridium thermocatricum</i>	75	56	306
36	1	181	2	gi 2088739	(AF003141) strong similarity to the FABY/P2/CRBP/CRABP family of transporters [Caenorhabditis elegans]	75	43	180
38	22	14510	15379	gi 1574058	[hypothetical [Haemophilus influenzae]	75	56	870
48	33	23398	24066	gi 1930092	[outer membrane protein [Campylobacter jejuni]	75	56	669
51	1	2	319	gi 43985	[nlS-like gene [Lactobacillus delbrueckii]	75	55	318
51	10	8318	11683	gi 537192	CG Site No. 620: alternate gene names hs, hsp, har, rmx apparent frameshift in GenBank Accession Number X06545 [Escherichia coli]	75	50	3366
54	18	19566	20759	gi 668669	[orf2 gene product [Lactobacillus leichmannii]	75	58	1194
57	9	8448	7822	gi 230561	[orf68 [Escherichia coli]	75	50	627
65	14	8072	6356	gi 606241	[30S ribosomal subunit protein S14 [Escherichia coli]	75	64	285
70	4	3071	2472	gi 125617	[adenine phosphoribosyltransferase [Bacillus subtilis]	75	57	600
71	24	30399	29404	gi 1574330	[C4 dicarboxylate transport protein [Haemophilus influenzae]	75	57	996
73	2	910	455	gnl PID#e249656	[yntf [Bacillus subtilis]	75	57	456
79	1	1810	491	gi 1146219	[28.2% of identity to the <i>Escherichia coli</i> GTP-binding protein Era; putative [Bacillus subtilis]	75	59	1320
82	6	6360	6536	gi 1655715	[BzBD [Rhodobacter capsulatus]	75	55	177
83	6	1938	2975	gnl PID#e323529	[putative PksX protein [Bacillus subtilis]	75	56	1038
93	111	7368	5317	gi 38989	[methionyl-tRNA synthetase [Bacillus stearothermophilus]	75	58	2032
93	13	9409	8699	gi 159193	[glutamine transport ATP-binding protein Q [Methanococcus jannaschii]	75	54	711
95	1	1795	47	gnl PID#e323510	[YloY protein [Bacillus subtilis]	75	57	1749
103	1	262	1186	gnl PID#e265928	[unknown [Mycobacterium tuberculosis]	75	64	825
104	1	691	915	gi 460026	[repressor protein [Streptococcus pneumoniae]	75	54	225
113	5	2951	3883	gnl PID#d101119	[ABC transporter subunit [Synechocystis sp.]	75	55	933
121	1	320	1390	gi 214531	[repressor of class I heat shock gene expression HrcA [Streptococcus mutans]	75	58	1071
127	6	2614	3000	gi 150051	[M. jannaschii predicted coding region MJ1558 [Methanococcus jannaschii]	75	44	387
137	18	10082	10687	gi 193116	[Glycoprotein S [Entamoeba histolytica]	75	52	606
149	111	8499	9338	gnl PID#d100582	[unknown [Bacillus subtilis]	75	55	840

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
151	6	9100	7673	[gi 40467]	[HaS polypeptide, part of C18 family [Citrobacter freundii]	75	57	1428
158	1	986	3	[gnl PID e253891]	[UDP-glucose 4-epimerase [Bacillus subtilis]	75	63	984
172	8	5653	6774	[gi 142978]	[glycerol dehydrogenase [Bacillus stearothermophilus]	75	56	1122
172	9	7139	9730	[gnl PID e268056]	[Unknown [Mycobacterium tuberculosis]]	75	58	2592
173	1	261	79	[gnl PID e236169]	[C1d5_6 [Caenorhabditis elegans]	75	50	183
185	3	3066	2014	[gi 1574806]	[spermidine/putrescine transport ATP-binding protein (potA) [Haemophilus influenzae]]	75	56	1053
191	6	5235	4213	[gi 149518]	[phosphoribosyl antranilate transferase [Lactococcus lactis]]	75	61	1023
226	2	1774	1181	[gi 2314588]	[A000642] conserved hypothetical protein [Helicobacter pylori]	75	65	594
231	1	1	153	[gi 40173]	[homolog of E. coli ribosomal protein L21 [Bacillus subtilis]]	75	57	153
234	1	2	418	[gi 2293259]	[A0008220] YcgI [Bacillus subtilis]	75	59	417
279	1	532	151	[gi 1119198]	[Unknown protein [Bacillus subtilis]]	75	50	402
291	7	3558	3827	[gi 40011]	[ORF17 (AA 1-161) [Bacillus subtilis]]	75	48	270
375	2	137	628	[gi 410137]	[ORR13 [Bacillus subtilis]]	75	58	492
6	120	16721	17560	[gi 2393323]	[A0008220] YcdI [Bacillus subtilis]	74	53	840
7	6	4682	6052	[gi 1354211]	[PEP12-like protein [Bacillus subtilis]]	74	50	1371
18	4	3341	2427	[gnl PID d101319]	[YcgI [Bacillus subtilis]]	74	54	915
21	6	5885	4800	[gi 1072381]	[glutamyl-aminopeptidase [Lactococcus lactis]]	74	59	1086
24	2	739	548	[gi 2314732]	[A000655] ABC transporter, permease protein (YaaE) [Helicobacter pylori]	74	46	192
25	1	2	367	[gnl PID d100932]	[H2O-forming NADH oxidase [Streptococcus mutans]]	74	63	366
38	18	11432	12964	[gi 537034]	[ORF_0488 [Escherichia coli]]	74	57	1593
48	10	8924	6669	[gi 1513069]	[P-type adenosine triphosphatase [Listeria monocytogenes]]	74	53	2256
55	11	11564	11401	[gnl PID e283110]	[fend [Staphylococcus aureus]]	74	64	564
61	2	1782	427	[gi 2393216]	[A0008220] putative UDP-N-acetyl muramate-alanine ligase [Bacillus subtilis]	74	55	1356
76	10	9914	8065	[gnl PID d101325]	[YqIB [Bacillus subtilis]]	74	54	1350
83	2	666	926	[pir C33486 C334]	[hscC homolog - Bacillus subtilis]	74	55	261
86	9	8385	8080	[gi 683585]	[prophenate dehydratase [Lactococcus lactis]]	74	55	906

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
102	5	5005	5652	[gi 143394	[OMP-PRPP transferase [Bacillus subtilis]	74	57	648
103	5	4364	3267	[gnl PID a23524	[YhN protein [Bacillus subtilis]	74	62	1098
108	7	6864	7592	[gnl PID e257631	[methyltransferase [Lactococcus lactis]	74	56	729
131	2	478	146	[gnl PID d101320	[YqgZ [Bacillus subtilis]	74	45	333
133	2	1380	919	[gnl PID a113025	[hypothetical protein [Bacillus subtilis]	74	60	462
137	9	6167	6787	[gnl PID d100479	[Na ⁺ -ATPase subunit D [Enterococcus hirae]	74	53	621
149	4	3008	3883	[gnl PID d100581	[high level kanamycin resistance [Bacillus subtilis]	74	55	876
157	2	243	824	[gi 1573373	[methylated-DNA-protein-cysteine methyltransferase (dat) [Haemophilus influenzae]	74	48	582
164	6	3515	4249	[gi 410131	[ORX7 [Bacillus subtilis]	74	48	735
167	7	5446	5201	[gi 413927	[ipa-3r gene product [Bacillus subtilis]	74	55	246
171	1	1	1818	[gnl PID d102251	[beta-galactosidase [Bacillus circulans]	74	62	1818
172	4	1064	2392	[gi 466474	[callobiose phosphotransferase enzyme II' [Bacillus stearothermophilus]	74	50	1329
185	1	326	3	[gi 1573646	[Mg(2+)-transport ATPase protein C (mgcC) (SP:P222037) [Haemophilus influenzae]	74	68	324
188	2	1089	2018	[gi 1573008	[ATP dependent translocator homolog (mba) [Haemophilus influenzae]	74	44	910
189	11	6491	7174	[gi 1661199	[sakacin A production response regulator [Streptococcus mutans]	74	60	684
210	1	520	1287	[gi 2293107	[UF0002201 YcmQ [Bacillus subtilis]	74	60	768
261	1	816	192	[gi 666983	[putative ATP binding subunit [Bacillus subtilis]	74	55	645
263	3	1619	3655	[gi 1663232	[similarity with <i>S. cerevisiae</i> hypothetical 137.7 kD protein in subtelomeric	74	42	2037
265	2	644	1227	[gi 49272	[Y repeat region [Saccharomyces cerevisiae]	73	57	1437
368	1	1	942	[gi 603988	[Apparaginase [Bacillus licheniformis]	74	64	384
7	16	113357	11921	[gnl PID d101324	[unknown [Saccharomyces cerevisiae]	74	39	942
17	10	5706	5449	[gnl PID e105362	[single strand DNA binding protein [Bacillus subtilis]	73	58	528
34	15	10281	9790	[gnl PID d102151	[ORF42c [Chlorella vulgaris]	73	46	492

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins: similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
40	12	9876	9226	[gi 1173517]	riboflavin synthase alpha subunit [Actinobacillus pleuropneumoniae]	73	55	651
55	2	3592	839	[gnl PID d101887]	cation-transporting Atfase PacI [Synechocystis sp. 1]	73	60	2754
55	18	17494	16586	[gnl PID e1265580]	unknown [Mycobacterium tuberculosis]	73	52	909
65	16	7213	7767	[gi 143419]	ribosomal protein L6 [Bacillus stearothermophilus]	73	60	555
66	3	3300	1659	[gnl PID e2659883]	LacF [Lactobacillus casei]	73	52	360
70	10	5557	5733	[gi 83761]	envelope protein [Human immunodeficiency virus type 1]	73	60	177
71	4	6133	8262	[gnl PID e122063]	ss-1,4-galactosyltransferase [Streptococcus pneumoniae]	73	45	2130
72	1	3	851	[gi 2293177]	[AF0002201] transporter [Bacillus subtilis]	73	50	849
76	7	7019	6195	[gnl PID d101325]	XgF [Bacillus subtilis]	73	66	825
76	12	10009	9533	[gi 1573066]	uridine kinase [uridine monophosphokinase] (udk) [Haemophilus influenzae]	73	54	477
80	7	8113	9372	[gi 1377833]	aminopeptidase [Bacillus subtilis]	73	60	1260
97	5	3389	1668	[gnl PID d101954]	dihydroxyacid dehydratase [Synechocystis sp. 1]	73	54	1722
98	9	6912	7619	[gnl PID a314991]	FTSE [Mycobacterium tuberculosis]	73	54	708
108	11	10328	10440	[gi 1388109]	regulatory protein [Enterococcus faecalis]	73	54	489
128	6	3632	4222	[gi 1685111]	lrtf091 [Streptococcus thermophilus]	73	63	591
138	2	1575	394	[gi 147326]	transport protein [Escherichia coli]	73	60	1182
140	13	12938	11903	[plr ES33402 E5334]	serine O-acetyltransferase (EC 2.3.1.30) - Bacillus stearothermophilus	73	55	636
162	5	5701	4991	[gnl PID e123511]	putative IhaQ protein [Bacillus subtilis]	73	50	711
164	4	2323	2790	[gi 1592076]	hypothetical protein (SP_P2576B) [Methanococcus jannaschii]	73	52	468
164	8	4815	5546	[gi 410137]	[ORF]X3 [Bacillus subtilis]	73	56	732
170	5	4394	5102	[gnl PID d100559]	homologue of unidentified protein of E. coli [Bacillus subtilis]	73	46	909
178	7	3893	4855	[gi 46242]	Involulation protein B, 5'-end [Rhizobium loti]	73	56	963
204	6	5056	4278	[gnl PID e214719]	PICR protein [Bacillus thuringiensis]	73	41	819
213	2	832	2037	[gi 1565296]	ribosomal protein S1 homolog; sequence specific DNA-binding protein [Leuconostoc lactis]	73	55	1206
231	2	84	287	[gi 40173]	homolog of E. coli ribosomal protein L21 [Bacillus subtilis]	73	61	204
237	1	2	505	[gi 1773151]	adenine phosphoribosyltransferase [Escherichia coli]	73	51	504

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
269	1	2	691	gnl PID 0101328	[Yqix [Bacillus subtilis]]	73	36	690
289	2	1272	832	lpir A02771 R7MC	[ribosomal protein L7/L12 - Micrococcus luteus]	73	66	441
343	1	14	484	gi 178825	[NE0002761 hypothetical 30.4 kD protein in manZ-cspC intergenic region [Escherichia coli]]	73	47	471
356	1	222	4	gi 2149905	[D-glutamic acid adding enzyme [Enterococcus faecalis]]	73	50	219
7	5	3165	4691	gnl PID 0101833	[amidase [Synchocystis sp.]]	72	52	1527
7	17	11743	13300	gnl PID e289141	[similar to hydroxymyristoyl-(acyl carrier protein) dehydratase [Bacillus subtilis]]	72	54	453
22	119	15637	16224	gnl PID 0101929	[ribosome releasing factor [Synchocystis sp.]]	72	51	588
33	17	12111	11425	gnl PID 0101190	[Orf3 [Streptococcus mutans]]	72	55	687
34	7	7147	5627	gi 396501	[aspartyl-tRNA synthetase [Thermus thermophilus]]	72	52	1521
38	23	15372	16085	pir H64108 H641	[L-ribulose-phosphate 4-epimerase (araD) homolog - Haemophilus influenzae [strain Rd KW20]]	72	54	714
39	5	5094	6905	gnl PID e254877	[unknown [Mycobacterium tuberculosis]]	72	56	1812
40	6	4469	4636	gi 153362	[lactose repressor [Streptococcus mutans]]	72	58	168
48	29	21729	12444	gi 2314329	[inhibin beta-A subunit [Ovis aries]]	72	33	207
48				(NE0006231)	[glutamine ABC transporter, permease protein (glnP) [Helicobacter pylori]]	72	49	696
50	5	4529	3288	gi 1750108	[YrbA [Bacillus subtilis]]	72	54	1242
51	3	1044	2282	gi 2293210	[AF0082201 YcbJ [Bacillus subtilis]]	72	54	1239
52	113	11681	113938	gi 162521	[deoxyribodipyrimidine photolyase [Bacillus subtilis]]	72	45	258
55	1	841	35	gi 8822518	[ORF_0304: GTG start [Escherichia coli]]	72	59	807
75	5	2832	3191	gnl PID e209886	[mercuric resistance operon regulatory protein [Bacillus subtilis]]	72	44	360
76	6	6229	5771	gi 142450	[airC protein [Bacillus subtilis]]	72	53	459
79	5	5065	4592	gi 2293279	[AF0082201 YcgG [Bacillus subtilis]]	72	46	474
87	14	14726	12309	gnl PID e321502	[putative Prik protein [Bacillus subtilis]]	72	52	2418
91	1	444	662	gi 500631	[Mvo1 gene product [Saccharomyces cerevisiae]]	72	50	219
91	7	4516	4764	gi 329615	[skeletal muscle sodium channel alpha-subunit [Equus caballus]]	72	38	249

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
95	2	2004	1717	gi PID e323527	putative Asp23 protein [Bacillus subtilis]	72	40	288
109	1	1452	118	gi 163331	alkaline phosphatase regulatory protein [Bacillus subtilis]	72	52	1335
126	1	3	2192	gi PID a101831	glutamine-binding periplasmic protein [Synechocystis sp.]	72	46	2190
130	3	1735	2478	gi 2415396	laf0157751 carboxypeptidase [Bacillus subtilis]	72	53	744
137	6	2585	2929	gi 472922	l-v-type Na-ATPase [Enterococcus hitzei]	72	46	345
140	10	9601	9203	gi 49224	URF 4 [Synechococcus sp.]	72	48	399
146	5	1906	1247	gi PID e323945	hypothetical protein [Bacillus subtilis]	72	45	660
147	2	2084	1083	gi PID e325016	hypothetical protein [Bacillus subtilis]	72	56	1002
147	5	6156	5146	gi 472327	TPP-dependent acetoate dehydrogenase beta-subunit [Clostridium magnum]	72	56	1011
148	8	5381	6433	gi 974332	NAD(P)H-dependent dihydroxyacetone phosphate reductase [Bacillus subtilis]	72	54	1053
148	14	10256	9675	gi PID d101319	YcpN [Bacillus subtilis]	72	50	582
159	8	4005	4949	gi 1788770	(AE000330) o63; 24 pct identical (44 gaps) to 318 residues from penicillin-binding protein 4*; PBE_BACSU SW: P32959 (451 aa) [Escherichia coli]	72	43	945
172	10	5907	10620	gi 7533387	unknown [Saccharomyces cerevisiae]	72	55	714
220	3	2862	3602	gi 157475	hypothetical [Haemophilus influenzae]	72	50	741
267	1	3	449	gi 280513	f170 [Escherichia coli]	72	48	447
281	2	699	540	gi PID d100964	homologue of spartokinase 2 alpha and beta subunits LysC of B. subtilis [Bacillus subtilis]	72	45	360
290	1	1018	14	gi 474195	this ORF is homologous to a 40.0 kd hypothetical protein in the hrb3' region from E. coli, Accession Number X16000 [Mycoplasma-like organism]	72	54	1005
300	1	63	587	gi 746399	transcription elongation factor [Escherichia coli]	72	50	525
316	1	1326	4	gi 158127	protein kinase C [Drosophila melanogaster]	72	40	1323
342	1	227	3	gi PID d10164	unknown [Bacillus subtilis]	72	54	225
354	1	1	1005	gi PID d102048	C. thermocellum beta-glucosidase; P26208 (985) [Bacillus subtilis]	72	52	1005
6	10	8134	110467	gi PID e264229	unknown [Mycobacterium tuberculosis]	71	57	2314
7	20	16231	15464	gi 18046	3'-oxoacyl-1'-acyl-carrier protein reductase [Cuphea lanceolata]	71	52	768
15	1	1297	2	gi PID d100571	replicative DNA helicase [Bacillus subtilis]	71	51	1296
15	4	4435	3869	gi 499384	orf189 [Bacillus subtilis]	71	47	567

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
18	6	5120	4218	gnl PID d101318	YqgG [Bacillus subtilis]	71	51	903
29	1	1	540	g1 1773142	similar to the 20.2kd protein in TETB-ENOA region of B. subtilis [Escherichia coli]	71	56	540
38	20	113327	113330	g1 537036	ORF_0158 [Escherichia coli]	71	48	504
51	12	15015	12676	g1 149528	dipeptidyl peptidase IV [Lactococcus lactis]	71	55	2340
55	23	121040	20585	g1 2343285	[AF015453] surface located protein [Lactobacillus rhamnosus]	71	58	456
60	2	705	265	g1 580920	rodo (gtaA) polypeptide (AA 1-673) [Bacillus subtilis]	71	44	441
71	18	24679	26226	g1 580920	rodo (gtaA) polypeptide (AA 1-673) [Bacillus subtilis]	71	44	1548
71	25	30587	30360	g1 606028	ORF_0414; Genplot suggests frameshift near start but none found [Escherichia coli]	71	50	228
72	6	5239	6729	g1 580835	lysine decarboxylase [Bacillus subtilis]	71	48	1491
72	14	111991	12878	g1 624085	similar to rat beta-alanine synthetase encoded by GenBank Accession Number S77881; contains ATP/GTP binding motif [Paramaecium bursaria Chlorella virus 1]	71	54	888
73	11	7769	7033	g1 1906594	[PN] [Rattus norvegicus]	71	42	237
74	6	10385	8517	g1 157373	[poly-L-tRNA synthetase (proS) [Haemophilus influenzae]	71	52	1869
81	9	5772	6578	g1 147404	mannose permease subunit II-M-Man [Escherichia coli]	71	45	807
86	5	4602	1604	gnl PID e322063	ss-1,4-galactosyltransferase [Streptococcus pneumoniae]	71	53	999
105	4	3619	4707	g1 2323341	[AF014460] PeoP [Streptococcus mutans]	71	58	1089
106	13	113557	112955	g1 1519287	[LemA [Listeria monocytogenes]	71	48	603
114	2	1029	1979	g1 310303	[mosA [Rhizobium meliloti]]	71	55	951
122	2	564	1205	g1 1649037	glutamine transport ATP-binding protein GLNQ [Salmonella typhimurium]	71	50	642
132	5	9018	7063	gnl PID d102049	H. influenzae hypothetical ABC transporter; P44808 (974) [Bacillus subtilis]	71	51	1956
140	1	1141	227	g1 1673788	[AE000015] Mycoplasma pneumoniae, fructose-bisphosphate aldolase; similar to Swiss-Prot Accession Number P1243; [from B. subtilis [Mycoplasma pneumoniae]]	71	49	915
140	5	5635	4973	gnl PID d100964	homologue of hypothetical protein in a ribamycin synthesis gene cluster of Streptomyces hygroscopicus [Bacillus subtilis]	71	48	663
141	7	7369	7845	gnl PID d102005	[AB001488] FUNCTION UNKNOWN, SIMILAR PRODUCT IN E. COLI AND MYCOPLASMA PNEUMONIAE. [Bacillus subtilis]	71	51	477

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
193	1	1	165	gi 146912	[ribosomal protein L13 [Staphylococcus carnosus]]	71	59	165
194	3	2205	1594	gi 5335351	[CcdC [Bacillus subtilis]]	71	52	612
199	3	1510	1319	gi 2102574	[IAE000090] Y4PE [Rhizobium sp. NG8234]	71	45	192
208	2	2616	3752	gi 1787378	[IAE000211] Hypothetical protein in purB 5' region [Escherichia coli]	71	57	1137
209	2	2022	1141	gi 41432	[FepC gene product [Escherichia coli]]	71	46	882
210	5	1911	3071	gi 49316	[orfF2 gene product [Bacillus subtilis]]	71	45	1161
210	6	3069	3386	gi 580900	[orfF3 gene product [Bacillus subtilis]]	71	48	318
212	2	3561	1381	gi 557367	[ribonucleotide reductase R1 subunit [Mycobacterium tuberculosis]]	71	53	2881
233	3	2003	12920	gi n P10 d101320	[Yqgr [Bacillus subtilis]]	71	50	918
244	1	13	1053	gnl P10 d100964	[homologue of asparokinase 2 alpha and beta subunits lysC of <i>B. subtilis</i>]	71	55	1041
251	2	1008	1874	gi 755601	[Bacillus subtilis]	71	46	867
282	2	906	712	gi 1353874	[unknown [Rhodobacter capsulatus]]	71	46	195
312	4	2137	1565	gnl P10 d102245	[LAB005534] YxbF [Bacillus subtilis]	71	34	573
338	1	3	683	gi 1591045	[hypothetical protein (SP:P14166) [Methanococcus jannaschii]]	71	48	681
346	1	3	164	gi 1591234	[hypothetical protein (SP:P42297) [Methanococcus jannaschii]]	71	36	162
374	1	619	2	gi 1397526	[clumping factor [Staphylococcus aureus]]	71	23	618
377	1	688	2	gi 397526	[clumping factor [Staphylococcus aureus]]	71	23	687
3	8	7419	6938	gnl P10 d100946	[unknown [Bacillus subtilis]]	70	42	462
3	10	8395	9075	gnl P10 d255543	[putative iron dependant repressor [Staphylococcus epidermidis]]	70	46	681
7	14	11024	10554	gnl P10 d100290	[undefined open reading frame [Bacillus stearothermophilus]]	70	55	771
7	18	14213	13719	gnl P10 d101090	[biotin carboxyl carrier protein of acetyl-CoA carboxylase [Synechocystis sp.]	70	56	495
9	2	1057	287	gnl P10 d100581	[unknown [Bacillus subtilis]]	70	52	771
12	4	2610	1789	gnl P10 d101195	[yycJ [Bacillus subtilis]]	70	52	822
21	2	2886	1846	gi 2293447	[ATP00830] ATPase [Bacillus subtilis]	70	54	741
22	13	10555	11512	gi 11165295	[Ydr540cp [Saccharomyces cerevisiae]]	70	50	558
30	6	4315	2980	gi 39478	[ATP binding protein of transport ATPases [Bacillus firmus]]	70	51	316

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
31	1	370	113	[gi 6627292	single-stranded DNA binding protein [unidentified eubacterium]	70	36	258
33	15	10639	9521	[gi 1161219	homologous to D-amino acid dehydrogenase enzyme [Pseudomonas aeruginosa]	70	50	1119
38	6	3812	4312	[gi 2058547	ComYD [Streptococcus gordoni]	70	48	501
38	125	17986	18477	[gi 537033	[ORF_f356 [Escherichia coli]	70	58	492
40	13	11054	9846	[gi 1173516	riboflavin-specific diaminase [Actinobacillus pleuropneumoniae]	70	52	1209
42	2	722	1954	[gi 1146183	putative [Bacillus subtilis]	70	51	1233
43	3	2373	1612	[gi 1159493	[glutamine transport ATP-binding protein Q [Methanococcus jannaschii]	70	48	762
45	8	9197	8049	[gnl PID d102036	subunit of ADP-glucose pyrophosphorylase [Bacillus stearothermophilus]	70	54	1149
59	2	567	956	[gnl PID d100302	neopullulanase [Bacillus sp.]	70	42	390
60	3	1874	795	[gnl PID e776466	aminopeptidase P [Lactococcus lactis]	70	48	1080
61	4	5553	2437	[gnl PID e275074	[SNF [Bacillus cereus]	70	51	3117
61	7	7914	6802	[gi 1573037	cystathione gamma-synthase (metB) [Haemophilus influenzae]	70	52	1113
63	7	5372	7222	[gnl PID d100974	unknown [Bacillus subtilis]	70	54	1851
68	7	7126	6962	[gi 1261014	[emm18_1 gene product [Streptococcus pyogenes]	70	37	165
72	112	10081	10911	[gi 2313093	[AE000524] carboxymorospemidine decarboxylase (nsPC) [Helicobacter pylori]	70	56	831
75	10	7888	8124	[gi 1877423	[galactose-1-P-uridylyl transferase [Streptococcus mutans]	70	59	237
79	3	3424	2525	[gi 39881	[ORF_311 (AA 1-111) [Bacillus subtilis]	70	47	900
87	10	9169	7324	[gnl PID e223506	putative Pkn2 protein [Bacillus subtilis]	70	52	2046
96	14	10640	11788	[gi 1573209	[tRNA-guanine transglycosylase (tgt) [Haemophilus influenzae]	70	52	1149
113	2	574	1086	[A180 [Saccharomyces cerevisiae]		70	59	513
123	5	2901	3461	[gnl PID d100585	unknown [Bacillus subtilis]	70	45	561
125	5	4553	4282	[gnl PID e276474	[tRNA-guanine transglycosylase (tgt) [Haemophilus influenzae]	70	35	312
129	5	4500	3454	[gnl PID d101314	capacitative calcium entry channel 1 [Bos taurus]	70	47	1047
133	3	2508	1394	[gi 223312	[YtfE [Bacillus subtilis]	70	50	1215
135	1	420	652	[gnl PID a265510	[YorfE [Streptococcus pneumoniae]	70	47	243
137	3	438	932	[gi 472919	[v-type Na-ATPase [Enterococcus hirae]	70	57	495
138	1	440	3	[gi 147336	transmembrane protein [Escherichia coli]	70	42	438

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
140	16	18796	16364	gi 976441	[N5-methyltetrahydrofolate homocysteine methyltransferase [Saccharomyces cerevisiae]]	70	53	2433
167	10	8263	6695	gi 149535	[D-alanine activating enzyme [Lactobacillus casei]]	70	52	1569
204	4	3226	2747	gnl PID d102049	[E. coli hypothetical protein: P11805 (267) [Bacillus subtilis]]	70	51	480
207	3	2627	2869	gnl PID e309213	[rcGAP [Dictyostelium discoideum]]	70	45	243
282	3	1136	882	gi 1353674	[unknown [Rhodobacter capsulatus]]	70	50	235
6	21	17754	18453	gnl PID e233879	[hypothetical protein [Bacillus subtilis]]	69	44	900
6	22	18482	19471	gi 580883	[Ipa-88d gene product [Bacillus subtilis]]	69	53	990
22	6	4682	5824	gi 2209379	[(AF006720) Proj [Bacillus subtilis]]	69	48	1143
22	9	7992	8651	gnl PID d100580	[unknown [Bacillus subtilis]]	69	51	660
22	12	9871	10767	gnl PID d100581	[unknown [Bacillus subtilis]]	69	51	897
27	7	5857	5348	gnl PID d102012	[(AB001488) FUNCTION UNKNOWN: [Bacillus subtilis]]	69	28	510
36	10	7294	10116	gi 437916	[isoleucy]-tRNA synthetase [Staphylococcus aureus]	69	53	2823
38	1	2	1090	gi 141900	[alcohol dehydrogenase (EC 1.1.1.1) [Alcaligenes eutrophus]]	69	48	1089
40	14	11333	11944	gi 1575280	[Holliday junction DNA helicase (ruvA) [Haemophilus influenzae]]	69	44	612
40	15	11942	12517	gi 1573653	[DNA-3'-methyladenine glycosidase I (tagI) [Haemophilus influenzae]]	69	50	576
45	6	6947	5490	gi 580887	[starch (bacterial glycogen) synthase [Bacillus subtilis]]	69	47	1458
48	34	124932	124153	gnl PID e233870	[hypothetical protein [Bacillus subtilis]]	69	36	780
49	6	6183	6521	gi 196297	[similar to phosphotransferase system enzyme II [Escherichia coli]]	69	50	339
49	8	7586	8338	gi 396420	[similar to Alcaligenes eutrophus pKG1 D-ribulose-5'-phosphate 3 epimerase [Escherichia coli]]	69	49	753
55	6	8262	7013	gi 1146238	[poly(A) polymerase [Bacillus subtilis]]	69	50	1230
59	3	954	2333	gnl PID e113038	[hypothetical protein [Bacillus subtilis]]	69	54	1380
62	3	1170	1418	gnl PID d101915	[hypothetical protein [Synochocystis sp.]]	69	49	249
63	8	7298	7762	gi 1293017	[ORF3 (put.); putative [Lactococcus lactis]]	69	42	465
66	4	3657	5081	gi 153755	[phospho-beta-D-galactosidase (EC 3.2.1.85) [Lactococcus lactis cremoris]]	69	49	1425
66	5	5126	6629	gi 433809	[enzyme II [Streptococcus mutans]]	69	46	1704
71	6	10017	10664	gnl PID a122063	[ss-1,4-galactosyltransferase [Streptococcus pneumoniae]]	69	39	648

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
71	121	22730	27966	gnl PID d100649	[Drosophila melanogaster] DE-cadherin	69	30	237
77	1	1	237	gi 207870	[Lactococcus lactis] groES gene product	69	44	237
81	5	3622	4101	gi 1573605	[Haemophilus influenzae] fucose operon protein (fucU)	69	52	480
83	1	40	714	ipr C33496 C334	[hsc homolog - Bacillus subtilis]	69	46	675
83	16	115742	116335	gi 143372	[phosphoribosyl glycinamide formyltransferase (PUR-N) [Bacillus subtilis]]	69	46	594
85	2	1212	916	gi 194097	[IINN-response element binding factor 1 [Mus musculus]]	69	48	297
91	5	1678	4274	gi 1574712	[anaerobic ribonucleoside-triphosphate reductase activating protein (nrdG) [Haemophilus influenzae]]	69	44	597
98	5	1247	4032	gnl PID d100262	[LvvP protein [Salmonella typhimurium]]	69	51	786
108	5	4085	5056	gnl PID d257629	[transcription factor [Lactococcus lactis]]	69	49	972
126	3	3078	4568	gnl PID d101329	[YcfJ [Bacillus subtilis]]	69	49	1491
131	6	4121	2889	gnl PID d10114	[YceR [Bacillus subtilis]]	69	47	1233
136	2	1505	2299	gnl PID d100581	[unknown [Bacillus subtilis]]	69	47	795
149	5	3852	4763	gnl PID d3232525	[YqQ protein [Bacillus subtilis]]	69	50	912
149	12	9336	10655	gi 151571	[Homology with E. coli and P. aeruginosa lyaA gene; product of unknown function; putative (Pseudomonas syringae)]	69	52	1320
153	4	3191	3829	gi 171073	[BunQ [Bacillus subtilis]]	69	44	639
169	3	849	2124	gnl PID d100582	[temperature sensitive cell division [Bacillus subtilis]]	69	49	1476
180	1	566	3	gi 488339	[alpha-amylase [unidentified cloning vector]]	69	50	564
212	1	1196	231	gi 1393209	[ribonucleotide reductase R2-2 small subunit [Mycobacterium tuberculosis]]	69	53	566
226	1	2	661	ipr J0285 J022	[indulin-26 - soybean]	69	41	660
233	5	3249	4766	gi 472918	[v-type Na-ATPase [Enterococcus hirae]]	69	56	1518
235	3	660	1766	gi 148945	[methylase [Haemophilus influenzae]]	69	43	1107
243	2	865	2361	gnl PID d100225	[ORF5 [Barley yellow dwarf virus]]	69	69	1497
251	3	2899	1967	gi 2289231	[macrolide-efflux protein [Streptococcus agalactiae]]	69	51	933
310	1	1	282	gnl PID d322442	[peptide deformylase [Clostridium bifermentation]]	69	55	282
369	1	868	2	gi 397526	[clumping factor [Staphylococcus aureus]]	69	22	867
370	1	749	3	gi 397526	[clumping factor [Staphylococcus aureus]]	69	21	747

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
379	1	44	280	gnl PfD d100669	DE-cadherin [Drosophila melanogaster]	69	30	237
388	1	260	72	gi 1787524	(AE000225) hypothetical 32.7 kd protein in trpB-truR intergenic region	69	44	189
1	2	2006	3040	gnl PfD d101809	ABC transporter [Synechocystis sp.]	68	43	1035
12	5	3958	2600	gi 2182992	histidine kinase [Lactococcus lactis cremoris]	68	45	1359
15	2	1790	1311	pir s16974 r5BS	ribosomal protein L9 - Bacillus stearothermophilus	68	56	480
16	6	7353	5701	gi 1787041	(AE000184) 0510; This 510 aa orf is 33 put. identical (4 gaps) to 525 residues of an approx. 640 aa protein YHEIN SW: P44808 [Escherichia coli]	68	45	1653
17	12	6479	6805	gi 5531165	acetylcholinesterase [Homo sapiens]	68	68	327
20	13	11128	14505	gi 142700	P competence protein (tg start codon) (put.); putative [Bacillus subtilis]	68	40	378
22	32	24612	25397	gi 289262	cone ORF3 [Bacillus subtilis]	68	36	786
30	7	4548	4288	gi 3111388	ORF1 [Azotobacter caulinodans]	68	46	261
36	5	1911	4585	gi 1573041	hypothetical [Haemophilus influenzae]	68	54	675
46	6	5219	6040	gi 1790331	(NE000446) hypothetical 29.7 kd protein in ibpa-gyrB intergenic region [Escherichia coli]	68	47	812
54	10	6235	7086	gi 882519	ICG Site No. 29739 [Escherichia coli]	68	55	852
55	5	7069	5165	gnl PfD d101914	ABC transporter [Synechocystis sp.]	68	45	1305
71	3	6134	5613	gi 1573353	outer membrane integrity protein (oia) [Haemophilus influenzae]	68	50	522
71	10	15342	16613	gi 580866	ipa-12d gene product [Bacillus subtilis]	68	31	1272
71	12	17560	18792	gi 14073	[secY protein [Lactococcus lactis]	68	35	1233
71	17	12295	24703	gi 1762349	involved in protein export [Bacillus subtilis]	68	50	2409
73	16	10208	9729	gi 1353337	ltpBpase (Bacteriophage rit)	68	43	480
86	18	117198	116011	gi 413943	ipa-19d gene product [Bacillus subtilis]	68	53	1188
87	17	17491	15866	gi 150209	ORF 1 [Mycobacteria mycolides]	68	43	1626
89	6	5139	4334	gi 1498824	M. jannaschii predicted coding region M20062 [Methanococcus jannaschii]	68	40	786
89	11	8021	8242	gi 150974	4-oxalocrotonate tautomerase [Pseudomonas putida]	68	43	222
97	8	6755	5394	gi 2367358	(AE000491) hypothetical 52.9 kd protein in aidB-rpsF intergenic region [Escherichia coli]	68	41	1362

TABLE 2 *S. pneumoniae* - Putative coding regions of novel proteins' similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
98	3	1418	2308	[gnl PID a00261]	[<i>LiVA</i> protein (<i>Salmonella typhimurium</i>)]	68	40	891
99	13	16414	17280	[gi 455363]	[regulatory protein (<i>Streptococcus mutans</i>)]	68	50	867
115	3	5054	3693	[gi 466474]	[cellobiose phosphotransferase enzyme II', (<i>Bacillus stearothermophilus</i>)]	68	44	1362
124	7	3394	3221	[gnl PID a00702]	[cut14 protein (<i>Schizosaccharomyces pombe</i>)]	68	56	174
125	2	2923	1922	[gi 450566]	[transmembrane protein (<i>Bacillus subtilis</i>)]	68	50	1002
132	2	4858	2888	[gnl PID a01732]	[DNA ligase (<i>Synechocystis</i> sp.)]	68	52	1971
140	7	7765	7580	[gi 120711]	[Unknown (<i>Saccharomyces cerevisiae</i>)]	68	47	186
150	1	539	3	[gi 402390]	[ADP-ribosylarginine hydrolase (<i>Mus musculus</i>)]	68	59	537
164	1	58	867	[gnl PID e055114]	[glutamate racemase (<i>Bacillus subtilis</i>)]	68	49	810
164	2	819	1835	[gnl PID e055117]	[hypothetical protein (<i>Bacillus subtilis</i>)]	68	50	1017
169	7	3946	4104	[pir B94545 B565]	[hypothetical protein - <i>Lactococcus lactis</i> subsp. <i>lactis</i> plasmid pSL2]	68	40	159
170	4	4247	4396	[gi 30146]	[spore coat protein (<i>Bacillus subtilis</i>)]	68	52	150
171	8	6002	7054	[gi 38722]	[precursor (aa -20 to 38) (<i>Acinetobacter calcoaceticus</i>)]	68	54	1053
198	3	2473	1871	[gnl PID a313075]	[hypothetical protein (<i>Bacillus subtilis</i>)]	68	46	603
211	2	969	1802	[gi 1439528]	[EFIC-mn (<i>Lactobacillus curvatus</i>)]	68	45	834
214	8	4926	4231	[gnl PID d102049]	[H. influenzae hypothetical protein P439 0 (182) (<i>Bacillus subtilis</i>)]	68	50	696
217	6	4955	5170	[gnl PID e326966]	[similar to <i>B. vulgaris</i> CHS-associated mitochondrial ... (reverse transcriptase) (<i>Arabidopsis thaliana</i>)]	68	36	216
218	7	3930	4745	[gi 2293198]	[EF008220] YcpP (<i>Bacillus subtilis</i>)	68	49	777
220	6	4628	4338	[gnl PID e325791]	[<i>WJ000005</i>] orf1 (<i>Bacillus megaterium</i>)	68	38	816
236	1	746	108	[gi 410117]	[ORF13 (<i>Bacillus subtilis</i>)]	68	51	291
237	2	675	1451	[gi 396348]	[homoserine transsuccinylase (<i>Escherichia coli</i>)]	68	46	539
250	4	771	1229	[gi 310899]	[ORF2 (<i>Synechococcus</i> sp.)]	68	44	363
254	1	517	155	[gi 1787105]	(AE000189) 0648 was 0669; This 669 aa orf is 40 pct identical (1 gaps) to 217 residues of an approx. 232 aa protein YBA_HA1N SW: P42247	68	50	459
337	1	1	774	[gnl PID e261920]	[putative orf (<i>Bacillus subtilis</i>) (<i>Escherichia coli</i>)]	68	47	774
345	1	3	653	[gi 149513]	[thiolydlate synthase (EC 2.1.1.45) (<i>Lactococcus lactis</i>)]	68	61	651

TABLE 2 *S. pneumoniae* - Putative coding regions of novel proteins similar to known proteins

contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
386	2	417	4	gi 1573353	outer membrane integrity protein (colA) [Haemophilus influenzae]	68	51	414
2	4	5722	4697	gi 1592141	M. jannaschii predicted coding region MJ1507 [Methanococcus jannaschii]	67	26	1026
3	6	5397	4591	gi 12593175	[AF008220] signal transduction regulator [Bacillus subtilis]	67	44	807
5	2	2301	574	gi 2313385	[AB000547] para-aminobenzoate synthetase (pabB) [Helicobacter pylori]	67	48	1728
6	19	116063	16758	gi 413931	ipa-7d gene product [Bacillus subtilis]	67	41	696
22	8	7694	7897	gi 1528962	pyrrole-5-carboxylate reductase [Actinidia deliciosa]	67	51	804
29	10	8335	9072	gi 468745	gtcr gene product [Bacillus brevis]	67	41	738
31	3	1379	585	gi 2425123	[AF019886] Pk8 [Dictyostelium discoideum]	67	49	795
32	11	8849	10150	gi 42029	[ORF1] gene product [Escherichia coli]	67	47	1102
36	116	14830	15546	gi 1592142	[ABC transporter, probable ATP-binding subunit [Methanococcus jannaschii]	67	43	717
38	9	4958	5392	gi 214803	[T22B3.3] [Caenorhabditis elegans]	67	47	435
38	121	113775	14512	gi 537037	[ORF_0216] [Escherichia coli]	67	52	738
45	9	10428	9181	gi 551710	[branching enzyme (glgB) (EC 2.4.1.18) [Bacillus stearothermophilus]	67	51	1248
48	23	18344	17514	gi 413949	ipa-25d gene product [Bacillus subtilis]	67	50	831
50	2	1773	952	gi 1PID d101330	[yqdQ] [Bacillus subtilis]	67	55	822
53	1	431	3	gi 1574291	[fimbrial transcription regulation repressor (fliB) [Haemophilus influenzae]	67	40	429
55	13	112740	11946	gi 1PID d252590	[ORF YD037c] [Saccharomyces cerevisiae]	67	51	795
61	9	9210	8329	gi 1PID d264711	[ATP-binding cassette transporter A [Staphylococcus aureus]	67	50	882
71	2	5614	6117	gi 1197667	[vitellogenin [Anolis pulchellus]	67	36	504
81	7	4489	4983	gi 1142714	[phosphoenolpyruvate:mannose phosphotransferase element IIB [Lactobacillus curvatus]	67	42	495
97	3	986	1366	gi 1PID d102235	[ATP-binding cassette transporter A [Staphylococcus aureus]	67	43	381
102	1	2357	3214	gi 1276746	[acyl carrier protein [Porphyra purpurea]	67	37	258
86	8	8140	6809	gi 1144744	[PSR] [Enterococcus hirae]	67	45	1332
106	3	1109	1987	gi 148921	[LipCD protein [Haemophilus influenzae]	67	43	879
115	4	5982	5656	gi 835750	[putative cellobiose phosphotransferase enzyme III [Bacillus subtilis]	67	44	327

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
115	7	8421	8077	gi 466673	cellobiose phosphotransferase enzyme II' [Bacillus stearothermophilus]	67	51	345
127	13	8127	7021	gi 47326	transport protein [Escherichia coli]	67	45	1107
136	3	2215	2859	gnl P0100581	unknown [Bacillus subtilis]	67	49	645
140	21	23317	20906	gnl P0101912	[phenylalanyl]-tRNA synthetase [Synechocystis sp.]	67	43	2412
146	6	2894	1893	gi 218294	histidine kinase [Lactococcus lactis cremoris]	67	44	1002
151	8	11476	11117	gnl P0100085	orfF129 [Bacillus cereus]	67	48	360
160	10	7453	8646	gi 2281317	orfB; similar to a Streptococcus pneumoniae putative membrane protein encoded by GenBank Accession Number X99100; inactivation of the OrfB gene leads to UV-sensitivity and to decrease of homologous recombination (plasmidic test) [Lactococcus 1	67	46	1194
163	3	3093	4505	gnl P0101317	[YqfR [Bacillus subtilis]	67	47	1407
167	8	6704	5454	gi 1161933	[DltB [Lactobacillus casei]	67	45	1251
169	4	2322	2879	gnl P0101331	[YqfG [Bacillus subtilis]	67	41	558
171	111	7656	8184	gi 153881	pneumococcal surface protein A [Streptococcus pneumoniae]	67	50	729
188	3	1930	3723	gi 1543975	[AbcB [Thermoaerobacterium thermosulfurigenes]	67	46	1794
189	6	3599	3141	gnl P010325178	[Hypothetical protein [Bacillus subtilis]	67	52	459
205	3	1663	2221	gi 606073	[OrfF_0169 [Escherichia coli]	67	47	549
207	4	2896	3456	gi 227674	[DtxR/iron regulated lipoprotein precursor [Corynebacterium diphtheriae]	67	49	561
217	3	4086	3703	gi 895750	[putative cellobiose phosphotransferase enzyme III [Bacillus subtilis]	67	42	384
246	2	291	662	gi 1842438	unknown [Bacillus subtilis]	67	43	372
252	1	2	745	gi 2351768	[PspA [Streptococcus pneumoniae]	67	41	744
265	3	1134	1811	gi 2313847	[AE000585] L-asparaginase II (ansB) [Helicobacter pylori]	67	42	678
295	1	1	375	gi 227674	[DtxR/iron regulated lipoprotein precursor [Corynebacterium diphtheriae]	67	43	375
1	7	4898	5146	gnl P010255179	[unknown [Mycobacterium tuberculosis]	66	56	249
3	1	389	3	gnl P010269548	[Unknown [Bacillus subtilis]	66	48	387
3	20	19267	20805	gi 39956	[IIGLc [Bacillus subtilis]	66	50	1939
4	3	2545	2718	gi 1787864	[AE000228] phage shock protein C [Escherichia coli]	66	36	174
5	9	13197	142592	gi 157491	[Fimbrial transcription regulation repressor (p1B) [Haemophilus influenzae]	66	46	606

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
9	4	2872	1451	[gnl PID e265928	[unknown [Mycobacterium tuberculosis]	66	43	1422
12	2	1469	1200	[gi 520407	[orf2; GTG start codon [Bacillus thuringiensis]	66	42	270
15	112	10579	9897	[gi 2314738	[AE000531] translation elongation factor EF-TS [tsf] [Helicobacter pylori]	66	49	1083
16	2	1312	734	[gnl PID d102245	[AB005554] yxbF [Bacillus subtilis]	66	38	480
22	3	1372	1851	[gi 1480916	[signal Peptidase type II [Lactococcus lactis]	66	51	1269
22	7	5828	7096	[gnl PID e206261	[gamma-glutamyl phosphate reductase [Streptococcus thermophilus]	66	50	945
22	20	16194	117138	[gnl PID e281914	[Yirtl [Bacillus subtilis]	66	40	447
30	2	530	976	[gi 2314379	[AE000627] ABC transporter, ATP-binding protein (YhcG) [Helicobacter pylori]	66	49	786
32	1	199	984	[gi 312444	[ORF2 [Bacillus caldolyticus]	66	44	1119
33	13	8352	7234	[gi 1387979	44% identity over 102 residues with hypothetical protein from Synechocystis sp. accession D64006_CD; expression induced by environmental stress; some similarity to glycosyl transferases; two potential membrane-spanning helices [Bacillus subtilis]	66	39	951
34	6	5658	4708	[gnl PID e250724	[orf2 [Lactobacillus sakei]	66	48	219
34	14	9792	9574	[gi 1590997	[M. jannaschii Predicted coding region MJ0272 [Methanococcus jannaschii]	66	46	663
35	16	15163	14501	[gi 1773322	[Cap54 [Staphylococcus aureus]	66	35	804
36	9	6173	6976	[gi 15118680	[minicell-associated protein DivIVA [Bacillus subtilis]	66	43	429
36	11	10396	10824	[bbs 1553146	[insulin activator factor, INSAF [human, Pancreatic insulinoma, Peptide Partial, 744 aa [Homo sapiens]	66	50	1392
48	1	28	1119	[gnl PID e125204	[hypothetical protein [Bacillus subtilis]	66	40	303
48	7	3810	4112	[gi 2182574	[AB000080] Y4P8 [Rhizobium sp. NGR1234]	66	52	807
52	4	3595	2789	[gi 1388565	[major cell-binding factor [Campylobacter jejuni]	66	43	1587
54	3	2662	1076	[gnl PID d101831	[glutamine-binding periplasmic protein [Synechocystis sp.]	66	44	558
61	10	9740	9183	[gnl PID e154144	[mdr gene product [Staphylococcus aureus]	66	44	1101
72	13	110893	11993	[gi 2313129	[AE000526] H. pylori predicted coding region Hp0049 [Helicobacter pylori]	66	43	792
74	9	13267	12476	[gi 1573941	[hypothetical [Haemophilus influenzae]	66	48	867
75	1	2	868	[gi 1574631	[nicotinamide mononucleotide transporter (pmuc) [Haemophilus influenzae]	66	40	1029
75	7	5303	4275	[gi 41312	[put.. EBG repressor protein [Escherichia coli]	66	40	1029

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim		% ident	length (nt)
						match	gene		
82	7	6813	8123	[gnl]P10 d255128	[trigger factor (Bacillus subtilis)]	66	53	1311	
83	3	905	1219	[gnl]C33496 C334	[hsc homolog - Bacillus subtilis]	66	44	315	
86	10	9407	8925	[gnl]6833584	[shikimate kinase (Lactococcus lactis)]	66	41	483	
88	10	7001	6060	[gi 2098719]	[putative fimbrial-associated protein (Actinomyces naeslundii)]	66	52	942	
89	1	951	4	[gi 410118]	[OREX19 (Bacillus subtilis)]	66	41	948	
93	7	3661	2711	[gi 1187936]	[AB002601 f288; This 298 aa orf is 51 pct identical (5 gaps) to 297 residues of an approx. 304 aa protein YCSN_BACSU SW: R42972 (Escherichia coli)]	66	49	951	
104	3	1805	3049	[gi 1469784]	[putative cell division protein FtsW (Enterococcus hirae)]	66	48	1245	
106	14	113576	114553	[gi 140027]	[homologous to E. coli gldB (Bacillus subtilis)]	66	52	678	
107	3	965	1864	[gi 144858]	[ORF A (Clostridium perfringens)]	66	49	900	
112	7	5718	6593	[gi 1609332]	[ppRA (Haemophilus influenzae)]	66	43	876	
115	1	3	1302	[gi 1727367]	[HyRIP (Saccharomyces cerevisiae)]	66	56	300	
122	1	3	566	[gnl]P10 d101328	[qY1 (Bacillus subtilis)]	66	41	564	
126	8	111759	11046	[gnl]P10 d101163	[ORF3 (Bacillus subtilis)]	66	39	714	
128	111	8201	8431	[gi 726288]	[growth associated protein GAP-43 (Xenopus laevis)]	66	41	231	
131	8	4894	4508	[gi 4866661]	[trans related protein (Saccharomyces cerevisiae)]	66	48	663	
140	13	1236	2574	[gi 40056]	[phoP gene product (Bacillus subtilis)]	66	41	885	
140	115	116318	15434	[gi 1658189]	[5,10-methylenetetrahydrofolate reductase (Erwinia carotovora)]	66	42	291	
146	12	7926	7636	[gnl]P10 d101140	[transposase (Synchocystis sp.)]	66	48	984	
147	6	7137	6154	[gi 472326]	[TPP-dependent acetoate dehydrogenase alpha-subunit (Clostridium magnum)]	66	46	996	
149	6	4435	5430	[gnl]P10 d101887	[pentose-5-phosphate-3-epimerase (Synchocystis sp.)]	66	42	822	
149	13	10734	11575	[gi 42271]	[pyruvate formate-lyase activating enzyme (NA 1-246) (Escherichia coli)]	66	41	309	
186	4	2578	2270	[gnl]P10 d101199	[ORF11 (Enterococcus faecalis)]	66	38	213	
207	2	2340	2597	[gnl]P10 e32183	[envelope glycoprotein gp160 (Human immunodeficiency virus type 1)]	66	46	258	
210	7	3358	3678	[gi 49318]	[ORF4 gene product (Bacillus subtilis)]	66	46	321	
217	8	5143	5355	[gi 9538]	[thrombin receptor (Cricetulus longicaudatus)]	66	33	234	
220	4	3875	3642	[gi 466648]	[lalternate name ORF of L23635 (Escherichia coli)]	66	41	309	

TABLE 2 *S. pneumoniae* - Putative coding regions of novel proteins similar to known proteins

Contig ID	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
223	1	1070	138	[gnl PID e247187	[zinc finger protein (Bacteriophage phi8)	66	45	933
224	2	1864	2640	[gnl 176399	[putative ABC transporter subunit (Staphylococcus epidermidis)	66	41	777
243	1	3	872	[db51 AB000617_2	[YcaH (Bacillus subtilis)	66	45	870
268	2	891	568	[gi 1517210	[putative transposase (Streptococcus pyogenes)	66	60	324
322	1	2	643	[gi 149836	[Zn protease (Methanococcus jannaschii)	65	34	732
5	10	113909	113178	[gi 1574292	[hypothetical (Haemophilus influenzae)	65	48	726
6	11	10465	11190	[gi 142854	homologous to E. coli radC gene product and to unidentified protein from Staphylococcus aureus (Bacillus subtilis)	65	42	243
7	2	647	405	[pir C64146 C641	[hypothetical protein H10259 - Haemophilus influenzae (strain Rd KW20)	65	50	576
7	7	6246	6821	[gnl PID 01323	[Yahu (Bacillus subtilis)	65	54	477
10	2	1873	1397	[gi 163111	[ORF 1 (Streptococcus pneumoniae)	65	45	795
16	3	1428	2222	[gnl PID e325010	[hypothetical protein (Bacillus subtilis)	65	40	459
21	4	3815	3357	[gnl PID e34910	[hypothetical protein (Staphylococcus sciuri)	65	42	609
22	134	125776	26384	[gi 1123010	[CDXA (Actinobacillus pleuropneumoniae)	65	38	1359
43	2	1648	290	[gi 1044826	[F14B5.1 (Caenorhabditis elegans)	65	45	795
48	13	100622	10856	[gi 1573390	[hypothetical (Haemophilus influenzae)]	65	37	639
48	122	117521	16883	[gi 1573391	[hypothetical (Haemophilus influenzae)]	65	38	495
48	125	19027	18533	[gnl PID e64484	[YCR020c, len:215 (Saccharomyces cerevisiae)]	65	32	1479
49	3	3836	5334	[gi 1480429	[putative transcriptional regulator (Bacillus stearothermophilus)]	65	42	819
50	6	5337	4519	[gi 171963	[tRNA isopentenyl transferase (Saccharomyces cerevisiae)]	65	46	861
52	15	14726	15568	[gi 1498745	[M. jannaschii predicted coding region M0912 (Methanococcus jannaschii)]	65	42	783
59	7	3963	4745	[gi 1496514	[orf zeta (Streptococcus pyogenes)]	65	65	984
68	3	2500	3483	[gi 887824	[ORF 0310 (Escherichia coli)]	65	42	1248
69	3	2171	1077	[gnl PID e311453	[unknown (Bacillus subtilis)]	65	55	705
69	7	6029	5325	[gi 809660	[deoxyribose phosphate aldolase (Bacillus subtilis)]	65	39	84
71	5	8536	9783	[gi 1573224	[glycosyl transferase IgC (GP:U14554_4) (Haemophilus influenzae)]	65	42	1095
72	8	7664	8527	[gnl PID e267589	[unknown, highly similar to several spermidine synthases (Bacillus subtilis)]	65	39	84

TABLE 2
Intragenic non-coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
76	5	5773	4097	gnl P0 d101723	DNA REPAIR PROTEIN RECN (RECOMBINATION PROTEIN N) (Escherichia coli)	65	44	1677
76	9	8099	7875	gnl 11574276	exodeoxyribonuclease, small subunit (XseB) (Haemophilus influenzae)	65	38	225
84	2	2870	2152	gnl 2313188	conserved hypothetical protein (Helicobacter pylori)	65	41	519
86	115	14495	13407	gnl P0 d101800	[AE000532] conserved hypothetical synthase (Symeochocystis sp.]	65	44	1089
87	3	3706	2423	gnl 151259	[HMG-CoA reductase (EC 1.1.1.88) (Pseudomonas mevalonii)	65	51	1284
88	3	2425	2336	gnl 1095510	unknown [Lactococcus lactis]	65	30	312
89	2	1627	1007	gnl P0 d102008	[AB001488] SIMILAR TO ORF14 OF ENTEROCOCCUS FAECALIS TRANSPOSON TN916.	65	41	621
111	6	6635	6186	gnl P0 d1e246063	[NN23]nucleoside diphosphate kinase (Xenopus laevis)	65	50	450
116	1	3	1016	gnl P0 d101125	[Synechocystis sp.] queuosine biosynthesis protein QueA	65	44	1014
123	1	69	389	gnl 498839	[ORF2 (Clostridium perfringens)]	65	36	321
123	7	6522	7190	gnl 1575577	[DNA-binding response regulator (Thermotoga maritima)]	65	39	669
125	3	3821	2859	gnl P0 d1e257609	[sugar-binding transport protein (Aerococcus thermophilum)]	65	47	963
137	12	8015	7818	gnl 2182574	[AE000090] Y4pE (Rhizobium sp. NGRB234)	65	41	198
147	4	5021	3885	gnl 1472329	[dihydrolipoamide acetyltransferase (Clostridium magnum)]	65	47	1137
148	2	1053	1931	gnl P0 d101319	[YqgH (Bacillus subtilis)]	65	42	879
151	2	3212	4687	gnl 304897	[EcoE type I restriction modification enzyme M subunit (Escherichia coli)]	65	50	1476
156	2	730	437	gnl 310893	[membrane protein (Thielliera parva)]	65	47	294
164	7	4256	4837	gnl 4010132	[ORF8 (Bacillus subtilis)]	65	48	582
169	6	3192	3914	gnl 1552737	[similar to purine nucleoside phosphorylase (Escherichia coli)]	65	43	732
176	4	2951	2220	gnl P0 d139500	[oligopeptide binding lipoprotein (Streptococcus pneumoniae)]	65	40	657
195	4	4556	3900	gnl 1592142	[ABC transporter, probable ATP-binding subunit (Methanococcus jannaschii)]	65	51	1413
196	1	160	1572	gnl P0 d102004	[AB001488] PROBABLE UDP-N-ACETYLGLUCOSAMINYL-D-GLUTAMYL-2, 6-DIAMINOLIGASE (EC 6.3.2.151). (Bacillus subtilis)	65	37	1032
204	2	2246	1215	gnl 143156	[membrane bound protein (Bacillus subtilis)]	65	48	348
210	4	1544	1891	gnl 49315	[ORF1 gene product (Bacillus subtilis)]	65	42	903
242	2	1625	723	gnl 1787540	[AB000226] f249: This 249 aa orf is 32 pct identical (8 gops) to 244 residues of an approx. 272 aa protein AGAR_ECOLI SW: P429202 (Escherichia coli)	65	42	

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
284	1	1	900	[gnl 1559861]	[clyM (Plasmid PAD1)]	65	36	900
304	1	2	574	[gnl PID e290934]	[unknown (Mycobacterium tuberculosis)]	65	52	573
315	1	2	1483	[gnl 790694]	[mannuronan C-5-epimerase (Azotobacter vinelandii)]	65	57	1482
320	1	3	569	[gnl PID d102048]	[K. aerogenes histidine utilization repressor; P12380 (1991) DNA binding protein (Bacillus subtilis)]	65	46	567
358	1	1	309	[gnl PID e32508]	[YLOS protein (Bacillus subtilis)]	65	55	309
2	7	7571	6696	[gnl 1498753]	[nicotinate-nucleotide pyrophosphorylase (Rhodospirillum rubrum)]	64	47	876
6	6	5924	6802	[gnl PID d10111]	[methionine aminopeptidase (Synechocystis sp.)]	64	58	270
8	4	3417	3686	[gnl 1045935]	[DNA helicase II (Mycoplasma genitalium)]	64	46	561
11	4	3249	2689	[gnl PID e265329]	[Orf8 (Streptococcus pneumoniae)]	64	45	642
15	7	6504	7145	[ycr59c yig2]	[yig2 homolog (Bacillus subtilis)]	64	38	348
22	11	9548	9695	[gnl PID d100381]	[Unknown (Bacillus subtilis)]	64	44	672
22	30	122503	123174	[gnl PID e283260]	[comE ORF1 (Bacillus subtilis)]	1	64	30
26	7	14375	14199	[gnl 409386]	[bmrU (Bacillus subtilis)]	64	51	177
27	2	1510	1334	[gnl 40795]	[DdeI methylase (Desulfovibrio vulgaris)]	64	50	318
29	2	614	297	[gnl 2326168]	[type VII collagen (Mus musculus)]	64	50	354
35	2	368	721	[pir Jc151 Jc11]	[hypothetical 20 kDa protein (insertion sequence IS1111) - Agrobacterium tumefaciens (strain P022) plasmid Ti]	64	41	447
40	1	3	449	[gnl 46970]	[lipid gene product (Staphylococcus epidermidis)]	64	45	294
40	7	4683	4976	[gnl PID e325792]	[ADP-glucose kinase (Bacillus megaterium)]	64	40	1149
45	7	8068	6920	[gnl PID d102036]	[subunit of ADP-glucose pyrophosphorylase (Bacillus stearothermophilus)]	64	54	759
51	2	201	1059	[gnl 43985]	[lntS-like gene (Lactobacillus delbrueckii)]	64	46	3107
51	13	15251	18397	[gnl 2293260]	[AT008220) DNA-polymerase III alpha-chain (Bacillus subtilis)]	64	47	603
53	3	1157	555	[gnl 1574292]	[hypothetical Haemophilus influenzae]	64	51	2631
58	2	4236	1606	[gnl 1573826]	[lalanyl-tRNA synthetase (Haemophilus influenzae)]	64	42	1257
66	1	3	1259	[gnl 895749]	[putative cellobiose phosphotransferase enzyme II' (Bacillus subtilis)]	64	47	1344
68	5	5213	6556	[gnl 436865]	[ImaiA1 gene products (Bacillus stearothermophilus)]	64	52	408
69	6	5356	4949	[gnl PID d101316]	[cdd (Bacillus subtilis)]	64	47	408

TABLE 2 *S. pneumoniae* - Putative coding regions of novel proteins & similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
81	113	14016	14231	[gi 143175	[Bacillus subtilis]	64	50	1911
83	122	21851	22090	[gnl PID 010315	[Bacillus subtilis]	64	57	183
87	111	10046	9300	[gnl PID e323505	[putative Ptc1 protein [Bacillus subtilis]	64	43	747
98	9	5012	5106	[gnl PID e333800	[hypothetical protein [Escherichia coli]	64	38	675
105	1	2	1276	[gi 1657503	[similar to <i>S. aureus</i> mercury(II) reductase [Escherichia coli]	64	45	1275
113	7	5136	6410	[gnl PID 010119	[<i>Naftobacterium pharaonis</i>]	64	40	1032
119	1	2	1297	[gnl PID e320520	[hypothetical protein [Natronobacterium pharaonis]	64	50	552
123	3	1125	2156	[gnl PID e253284	[ORF Ydu244w [Saccharomyces cerevisiae]	64	52	759
124	5	2331	1780	[gnl PID d101684	[hypothetical protein [Synechocystis sp.]	64	42	150
129	4	3467	2709	[gnl PID d101314	[YqeU [Bacillus subtilis]	64	50	354
131	1	152	3	[gi 1377841	[Unknown [Bacillus subtilis] - <i>Agrobacterium tumefaciens</i> (strain P022) plasmid Ti insertion sequence ::S1111]	64	44	576
137	11	7196	7549	[pir UC151 JCL1	[hypothetical 20.3K protein (strain P022) plasmid Ti]	64	45	1083
139	3	3226	2651	[gi 2293301	[YqB [Bacillus subtilis]	64	46	1017
146	110	6730	5648	[gi 1322245	[Imeavonate pyrophosphate decarboxylase [Rattus norvegicus]]	64	28	354
147	1	2	1018	[gnl PID e137033	[Unknown gene product [Lactobacillus leichmannii]]	64	31	702
148	111	8430	8783	[gi 2130630	[dynamin-like protein [Homo sapiens]]	64	43	816
156	7	4313	3612	[gnl PID d102050	[Unknown gene product [Streptococcus pyogenes]]	64	40	939
157	4	1299	2114	[gnl PID d100892	[Unknown gene product [Streptococcus pyogenes]]	64	43	816
162	6	5880	6362	[gi 5177204	[ORF1 putative 42 kDa protein [Bacillus subtilis]]	64	39	693
164	13	9707	8769	[gnl PID d100964	[System permease protein [Bacillus subtilis]]	64	33	354
175	5	3906	4598	[gi 1534045	[V. anguillarum [Bacillus subtilis]]	64	46	657
189	10	6154	6507	[gi 581207	[Antiterminator [Bacillus subtilis]]	64	46	657
191	4	3519	2863	[gi 149520	[Lactobacillus plantarum]]	64	46	657

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
1	1	76	1140	[gnl P01 e293806	[0-acetyl]homoserine sulfhydrylase [Leptospira, meyeri]	64	47	1065
202	1	234	1571	[gnl 1573393	[collagenase (ptc)] [Haemophilus influenzae]	64	42	1338
224	1	291	647	[gnl 40174	[ORF X] [Bacillus subtilis]	64	43	357
231	3	709	1089	[pi JCL151 JC11	[hypothetical 20.3K protein (insertion sequence IS1131) - Agrobacterium tumefaciens (strain P022) plasmid Ti]	64	50	381
253	1	820	2	[gnl 1377832	[Unknown] [Bacillus subtilis]	64	31	819
265	1	1	660	[gnl 1590871	[collagenase [Methanococcus jannaschii]	64	48	660
297	1	1	21	[gnl 99251	[Ginp] [Saccharomyces cerevisiae]	64	41	243
328	1	263	870	[gnl 556885	[Unknown] [Bacillus subtilis]	63	48	633
5	4	8720	8098	[gnl 1573101	[hypothetical] [Haemophilus influenzae]	63	40	696
10	6	5178	4483	[gnl 1573101	[membrane protein [Bacillus acidopolulliticum]]	63	42	579
12	11	9324	9902	[gnl 806536	[Unknown] [Acetobacter xylinum]	63	40	291
15	10	8897	9187	[gnl 722319	[ATP-dependent nuclease (Bacillus subtilis)]	63	32	723
17	2	1031	109	[gnl P01 e217602	[Unknown] [Lactobacillus plantarum]	63	45	804
18	8	7778	6975	[gnl 1377843	[Unknown] [Bacillus subtilis]	63	46	2703
26	4	9780	7078	[gnl 42440	[Unknown] [Bacillus subtilis]	63	35	705
29	5	3488	4192	[gnl 1377829	[Unknown] [Bacillus subtilis]	63	45	843
34	11	8830	7988	[gnl P01 d01198	[ORF8] [Enterococcus faecalis]	63	39	312
35	3	1187	876	[gnl 722339	[Unknown] [Acetobacter xylinum]	63	41	819
48	15	12509	11691	[gnl 1573389	[hypothetical] [Haemophilus influenzae]	63	35	531
51	11	12719	12189	[gnl 142450	[ahc] protein [Bacillus subtilis]	63	41	1044
55	4	3979	5022	[gnl 1708640	[YeaB] [Bacillus subtilis]	63	44	1002
55	15	113669	114670	[gnl P01 e211502	[thioredoxin reductase [Bacillus subtilis]]	63	43	906
68	10	9242	8919	[sp P37686 Y1AY_	[HYPOTHETICAL 40.2 KD PROTEIN IN AVTA-SELB INTERGENIC REGION (F382)]	63	40	324
96	8	5838	6484	[gnl 032803	[lic-1 operon protein (licb)] [Haemophilus influenzae]	63	41	870
86	7	6554	5685	[gnl 1574382	[putative fimbrial-associated protein (actinomycetes nasslundi)]	63	36	1701
88	8	6095	5180	[gnl 2088719	[putative fimbrial-associated protein (Streptococcus pneumoniae)]	63	38	627
96	8	5838	6484	[gnl 032803	[orfgyrb gene product [Streptococcus pneumoniae]]	63	36	1002
100	1	240	1940	[gnl 7171	[fuccosidase [Dictyostelium discoideum]]	63	43	906

TABLE 2 *S. pneumoniae* - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
104	4	1063	5765	gi 144985	phosphoenolpyruvate carboxylase [Corynebacterium glutamicum]	63	46	2703
106	8	9189	8554	gi 531099	endonuclease III [Bacillus subtilis]	63	45	636
122	6	4104	4886	gnl PID 0101139	transposase [Synnechocystis sp.]	63	39	183
128	7	4517	5203	gnl PID 0101434	orf2 [Methanobacterium thermoautotrophicum]	63	50	687
137	4	9633	1547	gi 472920	v-type Na-ATPase [Enterococcus hirae]	63	27	585
142	7	4100	4585	gnl PID 0313025	hypothetical protein [Bacillus subtilis]	63	44	486
159	5	1741	2571	gi 1787043	(AE000184) f271: This 271 aa orf is 24 pct identical (16 gaps) to 265 residues of an approx. 272 aa protein YIDh_ECOLI SW: P05997 [Escherichia coli]	63	39	831
171	112	8693	14406	gnl PID 0249498	19A1 protease [Streptococcus sanguis]	63	48	5604
177	1	3	347	gi 1773150	hypothetical 14.8kd protein [Escherichia coli]	63	34	345
178	2	423	917	gi 1722339	unknown [Acetobacter xylinum]	63	41	495
178	3	794	1012	gi 1591582	cobalamin biosynthesis protein N [Methanococcus jannaschii]	63	36	219
195	1	1377	175	gnl PID 0224217	ftsQ [Enterococcus hirae]	63	33	1203
234	5	1739	1527	gi 1591582	cobalamin biosynthesis protein N [Methanococcus jannaschii]	63	36	213
249	1	81	257	gi 1000453	[trr] [Bacillus subtilis]	63	41	177
283	1	127	1347	gi 396486	ORF8 [Bacillus subtilis]	63	44	1221
293	3	2804	3466	gi 722339	unknown [Acetobacter xylinum]	63	37	663
311	1	905	486	gi 1187744	UDP-galactose 4-epimerase [Streptococcus mutans]	63	46	420
324	1	2	556	gi 1477741	histidine periplasmic binding protein P29 [Campylobacter jejuni]	63	36	555
365	1	219	13	gi 2252843	(AF013293) No definition line found [Arabidopsis thaliana]	63	33	207
382	1	88	378	gi 722339	unknown [Acetobacter xylinum]	63	40	291
385	3	364	158	gi 1477743	(AF013293) No definition line found [Arabidopsis thaliana]	63	33	207
2	1	2495	288	gnl PID 0325007	penicillin-binding protein [Bacillus subtilis]	62	42	2288
3	123	123374	14231	gnl PID 0254993	hypothetical protein [Bacillus subtilis]	62	35	858
6	16	14320	13193	gnl PID 0349614	nifS-like protein [Mycobacterium leprae]	62	37	1128
7	8	6819	7232	gnl PID 010324	Yohy [Bacillus subtilis]	62	32	414
7	119	15466	14207	gnl PID 0101804	[beta ketoacyl acyl] carrier protein synthase [Synnechocystis sp.]	62	43	1260

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
7	121	17155	16229	gnl PID e321514	putative FabD protein [Bacillus subtilis]	62	46	927
7	124	19526	18519	gi 127634	[beta-ketoacyl-ACP synthase III [Cuphea wrightii]	62	37	1008
7	11	9678	9328	gi 1573768	[A/G-specific adenine glycosylase (mutY) [Haemophilus influenzae]	62	43	1203
12	7	5904	4702	gi 1591587	[pantothenate metabolism flavoprotein [Methanococcus jannaschii]	62	33	762
12	9	8032	8793	gi 1591587	[pantothenate metabolism flavoprotein [Methanococcus jannaschii]	62	43	351
15	11	9678	9328	gi 12151 gi 1	[hypothetical 20.3k protein (insertion sequence IS1131) - Agrobacterium tumefaciens (strain P022) plasmid Ti]	62	43	168
17	4	2609	2442	gi 11591081	[H. jannaschii predicted coding region HJ0374 [Methanococcus jannaschii]	62	44	219
17	5	3053	2835	gi 1149570	[role in the expression of lactacin F, part of the lacF operon [Lactobacillus sp.]	62	43	912
22	10	8627	9538	gnl PID d10580	[similar to B. subtilis DnaH [Bacillus subtilis]	62	43	1179
30	3	865	2043	gi 12314379	[AE00627] ABC transporter, ATP-binding protein (yhcG) [Helicobacter pylori]	62	44	600
33	5	2235	1636	gi 413976	[ipaS2r gene product [Bacillus subtilis]	62	34	435
38	11	5689	6123	gi 148231	[6251 [Escherichia coli]	62	43	945
40	17	14272	13328	gnl PID d101904	[hypothetical protein [Synochocystis sp.]	62	41	309
42	1	3	311	gi 1146182	[putative [Bacillus subtilis]	62	43	2739
44	2	1267	4005	gi 1786952	[AE000176] 0.877, 100 pct. identical to the first 86 residues of the 100 aa hypothetical protein Fragment Y86B_ECOLI SH: P24746 [Escherichia coli]	62	32	429
48	12	9712	9304	gi 662920	[repressor protein [Enterococcus hirae]	62	44	1518
51	8	5664	7181	gnl PID e301153	[systKI methylase [Salmonella enterica]	62	41	693
52	3	2791	2099	gi 1183886	[integral membrane protein [Bacillus subtilis]	62	40	999
55	16	15702	14704	gnl PID e313028	[hypothetical protein [Bacillus subtilis]	62	32	567
59	6	3418	3984	gi 2065483	[unknown [Lactococcus lactis lactis]	62	28	189
63	5	4997	4809	gi 1497771	[plin gene inverting protein (PlvM) [Moraxella lacunata]	62	45	738
70	14	10002	10739	gi 1992977	[bpLG gene product [Bordetella pertussis]	62	35	552
71	13	18790	20382	gi 1280135	[coded for by C. elegans cDNA cm01e2; similar to malibiose carrier protein (thiomethygalactoside permease 11) [Caenorhabditis elegans]	62	38	1284
71	128	32217	32768	gnl PID d101312	[YqEG [Bacillus subtilis]	62	62	1593
74	7	11666	10383	gi 1532753	[hypothetical [Escherichia coli]	62	35	1284

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	start (nt)	stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
80	8	9370	9609	gnl PID d102002	[FUNCTION UNKNOWN. [Bacillus subtilis]]	62	46	240
97	10	90688	7041	9i 1882463	[protein-N(pi)-phosphohistidine-sugar phosphotransferase (Escherichia coli)]	62	42	2028
98	4	2306	3268	gnl PID d101496	[BraB (integral membrane protein) (Pseudomonas aeruginosa)]	62	42	963
102	3	2823	3539	gnl PID e313010	[hypothetical protein (Bacillus subtilis)]	62	24	717
103	3	2795	1242	gnl PID d102059	[H. influenzae hypothetical ABC transporter; P44808 (974) [Bacillus subtilis]]	62	41	1554
111	2	2035	3462	9i 1581297	[Misp [Lactococcus lactis]]	62	44	1428
112	4	3154	4080	9i 1574379	[lic-1 operon protein (licA) (Haemophilus influenzae)]	62	39	927
112	6	4939	5639	9i 1573381	[lic-1 operon protein (licC) (Haemophilus influenzae)]	62	39	711
124	3	1137	721	9i 1573024	[anaerobic ribonucleoside-triphosphate reductase (nrdB) (Haemophilus influenzae)]	62	45	417
124	6	3162	2349	9i 609076	[leucyl aminopeptidase (Lactobacillus delbrueckii)]	62	40	834
126	7	11073	7516	gnl PTD d101163	[ORF4 [Bacillus subtilis]]	62	38	3558
129	6	4983	4540	9i 5105915415	[zinc finger protein EF6 - Chilo iridescent virus]	62	48	444
131	7	4510	4103	9i 1857245	[unknown [Lactococcus lactis]]	62	42	408
149	2	1923	2579	9i 1592142	[ABC transporter, probable ATP-binding subunit [Methanococcus jannaschii]]	62	41	657
149	7	5360	6055	gnl PID e323508	[yios protein [Bacillus subtilis]]	62	40	696
156	1	450	238	gnl PID e254644	[membrane protein [Streptococcus pneumoniae]]	62	40	213
156	6	3606	2935	gnl PID d102050	[transmembrane [Bacillus subtilis]]	62	37	672
171	2	1779	2291	9i 143941	[EIII-B Sor PPS [Klebsiella pneumoniae]]	62	35	513
172	1	385	723	9i 1895750	[putative cellobiose phosphotransferase enzyme III [Bacillus subtilis]]	62	39	339
173	3	2599	893	9i 11591732	[cobalt transport ATP-binding protein O [Methanococcus jannaschii]]	62	42	1707
179	2	492	1754	9i 11574071	[H. influenzae predicted coding region H11038 [Haemophilus influenzae]]	62	42	852
181	6	2856	3707	9i 1777435	[Lact [Lactobacillus casei]]	62	41	1764
185	2	2074	311	9i 2182297	[AE000073] Y4IN [Rhizobium sp. NGR334]	62	37	924
200	2	1061	1984	9i 450566	[transmembrane protein [Bacillus subtilis]]	62	41	891
202	3	2583	3473	9i 42219	[P35 gene product (AA 1 - 314) [Escherichia coli]]	62	45	192
210	3	1374	1565	9i 49315	[ORF1 gene product [Bacillus subtilis]]	62	41	1764

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
211	1	3	971	[gi 147402	mannose permease subunit III-Man [Escherichia coli]	62	43	969
223	2	1495	1034	[gnl PID1d101190	[Streptococcus mutans]	62	41	462
228	1	34	909	[gi 530063	glyceral uptake facilitator [Streptococcus pneumoniae]	62	44	876
234	2	90	917	[gi 2293259	[AF008220] YQ1 [Bacillus subtilis]	62	38	828
282	5	1765	1487	[gnl PID1e276475	galactokinase [Arabidopsis thaliana]	62	33	279
375	1	1	159	[gi 1674231	[AE000052] Mycoplasma pneumoniae, hypothetical protein homolog, similar to P515, from B. subtilis [Mycoplasma pneumoniae]	62	40	159
385	5	584	357	[gi 1573353	outer membrane integrity protein (tolA) [Haemophilus influenzae]	62	47	228
3	19	18550	19269	[gi 606162	[ORF f229] [Escherichia coli]	61	41	720
7	4	2725	3225	[gi 2114425	similar to Synechocystis sp. hypothetical protein, encoded by GenBank Accession Number D6006 [Bacillus subtilis]	61	43	273
17	6	3326	3054	[gi 149569	[lactacin F [Lactobacillus sp.]	61	38	897
44	3	4061	4957	[gnl PID1d10168	xylose repressor [Synochocystis sp.]	61	42	1155
54	111	8388	7234	[gnl PID1d101329	[YqjH [Bacillus subtilis]	61	42	2064
57	6	3974	6037	[gnl PID1d101316	[YqfK [Bacillus subtilis]	61	34	792
58	5	7356	6565	[sp P45169 POTC_	ISPERMIDINE/PUTRESCINE TRANSPORT SYSTEM PERMEASE PROTEIN POTC.	61	46	690
67	1	3	692	[gi 537108	[ORF f254] [Escherichia coli]	61	41	927
68	9	8816	7890	[gi 19501	[PPLZ12] gene product (AA 1-184) [Lupinus polyphyllus]	61	44	1272
70	115	110737	12008	[gi 992976	[bpfF gene product [Bordetella pertussis]	61	36	444
72	111	9759	10202	[gnl PID1d101833	carboxynospermidine decarboxylase [Synochocystis sp.]	61	45	879
76	8	7881	7003	[gnl PID1d100305	[farneoyl diphosphate synthase [Bacillus stearothermophilus]	61	42	1218
87	4	4914	3697	[gi 528991	[unknown [Bacillus subtilis]	61	44	951
87	113	12311	111361	[gi 1789883	[AE000407] methionyl-tRNA formyltransferase [Escherichia coli]	61	36	1491
115	6	7968	6478	[gi 895747	[putative cel operon regulator [Bacillus subtilis]	61	40	1338
123	8	7181	8518	[gi 11209527	[protein histidine kinase [Enterococcus faecalis]	61	44	789

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
126	6	7525	6725	gi 1787043	(AE000184) t271; This 271 aa orf is 24 pct identical (16 gaps) to 265 residues of an approx. 272 aa protein YIDA_ECOLI SW: P09397 [Escherichia coli]	61	38	801
128	1	1	639	gnl PID d101328	[Yq1Y (Bacillus subtilis)]	61	41	639
139	7	4794	5054	gi 1022726	Unknown [Staphylococcus haemolyticus]	61	41	261
139	9	12632	5913	gnl PID e270014	beta galactosidase [Thermoaerobacter ethanolicus]	61	41	6720
143	1	2552	42	gi 520541	penicillin-binding protein 1A and 1B [Bacillus subtilis]	61	42	2511
148	16	12125	11424	gi 1552743	tetrahydrodipicolinate N-succinyltransferase [Escherichia coli]	61	42	702
162	3	4112	3456	gnl PID d101829	phosphoglycolate phosphatase [Synechocystis sp.]	61	30	657
172	3	727	1077	gnl PID d102048	[B. subtilis, cellobiose phosphotransferase system, celA; P46318 (220) Bacillus subtilis]	61	44	351
177	3	1101	1772	gnl PID d100574	Unknown [Bacillus subtilis]	61	43	672
202	2	1278	2585	gi 1045831	hypothetical protein (GB:U1895_6) [Mycoplasma genitalium]	61	36	1308
224	3	2782	3144	gi 1591144	[M. jannaschii predicted coding region M0440] [Methanococcus jannaschii]	61	30	363
225	4	3395	3766	gi 1552774	hypothetical [Escherichia coli]	61	40	372
249	2	212	802	gi 1000453	[TrER (Bacillus subtilis)]	61	42	591
254	2	843	484	gnl PID d100417	ORF120 [Escherichia coli]	61	36	360
257	1	3	350	gnl PID e255315	Unknown [Mycobacterium tuberculosis]	61	42	348
293	4	3971	3657	pir JC1151JC11	hypothetical 20 kDa protein (insertion sequence IS1111) - Agrobacterium tumefaciens (strain P022) plasmid Ti	61	45	315
301	1	949	17	gi 12291209	[AF016424] contains similarity to acyltransferases [Caenorhabditis elegans]	61	33	933
373	1	1066	287	gi 393396	Tb-228 membrane associated protein [Trypanosoma brucei subgroup]	61	38	780
3	24	24473	124955	gi 537093	[ORF_0153b [Escherichia coli]]	60	27	483
6	5	4636	5739	gi 12293258	[AF008220] Yto1 [Bacillus subtilis]	60	35	1104
6	12	11936	11187	gi 293017	ORF3 (put.) : putative [Lactococcus lactis]	60	44	750
17	13	6708	6484	gi 149569	[lactacin F [Lactococcus sp.]	60	32	225
18	7	6977	5670	gi 1788140	(AE000278) o481; This 481 aa orf is 35 pct identical (19 gaps) to 309 residues of an approx. 856 aa protein NOLI_HUMAN SW: P46087 [Escherichia coli]	60	43	1108
20	15	15878	17167	gnl PID d100584	Unknown [Bacillus subtilis]	60	44	1290

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig ID	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
22	1	1	243	[gnl PID d102050	transmembrane [Bacillus subtilis]	60	36	243
32	110	8296	8964	[gnl 2293275	[TAFO08220] Ytag [Bacillus subtilis]	60	37	669
38	15	8837	9697	[gnl 40023	[B. subtilis genes rpmH, rpmB, 50kd, gida and qidB [Bacillus subtilis]	60	35	861
43	6	8610	5944	[gnl 171787	[protein kinase 1 [Saccharomyces cerevisiae]	60	36	2667
44	1	1	1269	[gnl PID e235823	[unknown [Schizosaccharomyces pombe]	60	44	1269
45	110	11138	10368	[gnl 397488	[1,4-alpha-glucan branching enzyme [Bacillus subtilis]	60	43	771
48	119	15766	14378	[gnl PID e203173	[orf1 [Lactobacillus helveticus]	60	39	1389
48	121	16727	16951	[gnl PID d102041	[AB002668] unnamed protein product [Haemophilus actinomycetemcomitans]	60	32	225
50	1	2	898	[gnl PID e248537	[ORP286 protein [Pseudomonas stutzeri]	60	31	897
62	2	618	1177	[gnl PID d102087	[unknown [Bacillus subtilis]	60	42	540
68	4	3590	5203	[gnl 1573583	[H. influenzae predicted coding region H10594 [Haemophilus influenzae]	60	36	1614
70	11	5781	6182	[gnl PID d102014	[AB00188] SIMILAR TO YDFR GENE PRODUCT OF THIS ENTRY (YDFR-BACSU) [Bacillus subtilis]	60	33	402
70	12	6343	8133	[gnl PID e124910	[hypothetical protein [Bacillus subtilis]	60	33	2457
71	8	111701	14157	[gnl 580866	[ipa-12d gene product [Bacillus subtilis]	60	45	846
74	8	12509	11664	[gnl PID d101822	[phosphatidate cytidyltransferase [Synchocystis sp.] [Bacillus subtilis]	60	38	1791
76	4	4116	3367	[gnl 2352096	[orf; similar to serine/threonine protein phosphatase [Fervidobacterium islandicum]	60	30	294
80	4	7372	7665	[gnl 1786420	[AE000131] f86; 100 pct identical to GB: ECODINJ_6 ACCESSION: D36582 [Escherichia coli]	60	39	750
81	6	4073	4522	[gnl 147402	[mannose permease subunit III-Man [Escherichia coli]	60	35	450
86	1	940	155	[gnl 143177	[putative [Bacillus subtilis]	60	26	786
92	1	1	192	[gnl 396348	[homoserine transsuccinylase [Escherichia coli]	60	45	192
93	14	10619	9384	[gnl 1788389	[AE000297] of64; This 464 aa orf is 31 pct identical (9 gaps) to 311 residues of an approx. 416 aa protein MTBC_NEIGO SW: P43505 [Escherichia coli]	60	27	1236
94	5	5548	8121	[gnl PID e329895	[AJ000496] cyclic nucleotide-gated channel beta subunit [Rattus norvegicus]	60	50	2574
97	7	5396	4533	[gnl 1591396	[Methanococcus jannaschii] transketolase	60	43	864
102	2	2081	2833	[gnl PID e202929	[hypothetical protein [Mycobacterium tuberculosis]]	60	43	753

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF	start	Stop	match	match	% sim	% ident	length
ID	ID	(nt)	(nt)	accession	gene name			(nt)
106	9	9773	9183	[gnl PID e334782]	[Y1BN protein [Bacillus subtilis]	60	31	591
113	8	6361	6837	[gi 466875]	[nitu; B1496_C1_157 (Mycobacterium leprae)]	60	43	477
115	2	2755	524	[gnl PID e228143]	[AD000332; Glucosidase II [Homo sapiens]]	60	32	2232
122	7	4763	5068	[gnl PID d01876]	[transposase [Synechocystis sp. 1]]	60	39	306
127	8	4510	5283	[gi 1777938]	[Pgm [Treponema pallidum]]	60	38	774
138	4	3082	2672	[gnl PID e225196]	[hypothetical protein [Bacillus subtilis]]	60	36	411
139	1	177	4	[gnl PID d00680]	[ORF [Thermus thermophilus]]	60	39	174
139	11	14520	13009	[gi 537145]	[ORF f437 [Escherichia coli]]	60	30	1512
140	12	12592	1249	[gi 1209527]	[protein histidine kinase [Enterococcus faecalis]]	60	37	1344
141	1	210	1049	[gi 463181]	[ES ORF from bp 3842 to 4081; putative [Human Papillomavirus type 33]]	60	34	840
141	5	5368	6405	[gi 145362]	[tyrosine-sensitive DAMP synthase (arof) [Escherichia coli]]	60	41	1038
142	6	3558	4049	[gi 600711]	[putative [Bacillus subtilis]]	60	37	492
148	10	7742	8713	[gnl PID e113022]	[hypothetical protein [Bacillus subtilis]]	60	27	972
153	5	3667	4278	[gi 1223322]	[IAF08220; branch-chain amino acid transporter [Bacillus subtilis]]	60	42	612
155	1	1413	748	[gi 12104504]	[putative UDP-glucose dehydrogenase [Escherichia coli]]	60	40	666
158	3	3116	2472	[gnl PID d00872]	[a negative regulator of rho regulon [Pseudomonas aeruginosa]]	60	37	645
159	3	778	1386	[gnl PID e308090]	[product highly similar to Bacillus anthracis CapA protein [Bacillus subtilis]]	60	48	609
163	7	8049	8468	[gnl PID d01313]	[Ygen [Bacillus subtilis]]	60	38	420
170	3	4130	2668	[gi 15747179]	[H. influenzae predicted coding region H1244 [Haemophilus influenzae]]	60	39	1443
171	7	4717	5901	[gi 606076]	[ORF_o384 [Escherichia coli]]	60	44	1185
183	3	2440	2135	[gi 1877427]	[repressor [Streptococcus pyogenes phage T12]]	60	38	306
191	10	9444	8428	[gi 415664]	[catabolite control protein [Bacillus megaterium]]	60	42	1017
200	1	139	1083	[gi 418462]	[transmembrane protein [Bacillus subtilis]]	60	37	945
201	3	3895	1928	[gi 475112]	[enzyme IIabc [Pediococcus pentosaceus]]	60	39	1968
214	15	10930	10439	[gi 1573407]	[hypothetical [Haemophilus influenzae]]	60	39	492
218	4	2145	2363	[gi 608320]	[myosin heavy chain kinase A [Dictyostelium discoideum]]	60	31	219

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF	ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
226	4	2518	2351	1911437705	[gi 437705	hyaluronidase [Streptococcus pneumoniae]	60	53	168
242	1	725	3	191143938	[gi 43938	Sor regulator [Klebsiella pneumoniae]	60	41	723
245	1	1	288	1911304897	[gi 304897	Eco type I restriction modification enzyme M subunit [Escherichia coli]	60	56	288
251	1	905	45	1911671632	[gi 671632	unknown [Staphylococcus aureus]	60	36	861
259	1	969	92	191153794	[gi 53794	rgg [Streptococcus gordonii]	60	32	888
260	2	1432	1682	1911318401	[gi 318401	S318 [probable transposase - Bacillus stearothermophilus	60	26	171
274	1	836	96	1911592173	[gi 592173	N-ethylammonium chlorohydrolase [Methanococcus jannaschii]	60	40	741
308	1	463	2	19111787397	[gi 1787397	[AE000214] o157 [Escherichia coli]	60	43	462
318	1	3	308	19119754	[gi 9754	[xerc recombinase [Lactobacillus leichmannii]	60	42	306
344	1	73	522	1911509672	[gi 509672	repressor protein [Bacteriophage Tuc2009]	60	32	450
5	1	576	4	19112293147	[gi 2293147	[AF008220] YtxM [Bacillus subtilis]	59	31	573
7	22	18140	17142	1911680724	[gi 680724	unknown [Mycobacterium tuberculosis]	59	39	999
10	1	1413	4	19111353880	[gi 1353880	sialidase L [Macrobdella decora]	59	41	1410
15	6	6463	5156	1911580841	[gi 580841	[F1 [Bacillus subtilis]	59	35	1308
22	2	479	1193	1911142469	[gi 142469	als operon regulatory protein [Bacillus subtilis]	59	34	915
22	5	2638	4614	191113280623	[gi 13280623	[PCPA [Streptococcus pneumoniae]	59	44	1917
30	1	208	558	1911433868	[gi 433868	[hypothetical protein [Bacillus subtilis]	59	37	351
30	4	3698	2455	1911102290	[gi 102290	unknown [Lactobacillus sake]	59	33	1224
35	13	12201	11071	19111071	[gi 1071	[hypothetical protein [Bacillus subtilis]	59	35	1131
35	14	13288	12182	19111657647	[gi 1657647	[CapBH [Staphylococcus aureus]	59	39	1107
36	18	18036	17897	19111500535	[gi 1500535	M_jannaschii predicted coding region M1615 [Methanococcus jannaschii]	59	33	180
38	12	6172	7137	1911229239	[gi 229239	[AF008220] YtxM [Bacillus subtilis]	59	40	1410
42	3	1932	3361	1911688845	[gi 688845	[pin1n [Canis familiaris]	59	41	951
50	3	2678	1728	1911101329	[gi 101329	[YqjK [Bacillus subtilis]	59	41	966
56	5	1870	2388	1911137594	[gi 137594	xerc recombinase [Lactobacillus leichmannii]	59	41	519
61	6	6812	5628	1911131516	[gi 131516	aminotransferase [Bacillus subtilis]	59	40	1185
67	5	2382	3023	1911146190	[gi 146190	[2-keto-3-deoxy-6-phosphogluconate aldolase [Bacillus subtilis]	59	36	642

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	orf ID	Start ID	Stop ID	match accession	match gene name	% sim	% ident	length (nt)
115	8	8766	8511	gi 1590886	[antohenate kinase (coA) [Haemophilus influenzae]	59	38	333
69	10	8567	8839	gi 1573628	[putative Fnu protein [Bacillus subtilis]	59	44	1329
87	12	11382	10055	gi PID a323504	[Mycoplasma pneumoniae, fructose- permease IIBC component; similar to Swiss-Prot Accession Number P20966, from E. coli [Mycoplasma pneumoniae]	59	43	1968
113	14	13927	15894	gi 1673733	[AE000010] Mycoplasma pneumoniae, fructose- permease IIBC component; similar to Swiss-Prot Accession Number P20966, from E. coli [Mycoplasma pneumoniae]	59	38	246
119	2	1966	1576	gi 1590886	[M. jannaschii predicted coding region N40110 [Methanococcus jannaschii]	59	43	441
128	17	13438	13178	gi 1PID e209005	[homologous to ORF2 in nrdEF operons of E.coli and S.typhimurium	59	38	261
140	22	23903	23388	gi 482922	[Lactococcus lactis]	59	40	516
148	13	9697	9014	gi 1PID d102005	[AB001488] FUNCTION UNKNOWN, SIMILAR PRODUCT: IN H. INFLUENZAE AND SYNCHOCYSTIS. [Bacillus subtilis]	59	32	684
149	10	7213	8244	gi 710432	[Cmp-binding factor 1 [Staphylococcus aureus]	59	40	1032
164	9	6993	6013	gi 1PID d100965	[ferric anguibactin-binding protein precursor FabB of V. anguillarum	59	41	981
164	12	8836	7823	gi 1PID d100964	[homologue of ferric anguibactin transport system permease protein FabC of V. anguillarum [Bacillus subtilis]	59	35	1014
177	2	401	1072	gi 289759	[coded for by C. elegans cDNA CE2G3 (GenBank:Z14728); putative Icanenorhabditis elegans]	59	40	672
177	7	3841	4200	gi 2313445	[AE000551] H. pylori predicted coding region HP0342 [Helicobacter pylori]	59	38	160
183	4	2768	2508	gi 1509672	[repressor protein [Bacteriophage Tuc2009]	59	50	261
186	6	3398	2820	gi 606080	[ORF_0290; Geneplot suggests frameshift linking to 0267, not found in Escherichia coli]	59	38	579
190	3	3120	1711	gi 1613768	[histidine protein kinase [Streptococcus pneumoniae]	59	32	1410
194	2	1621	1019	gi 1PID d100579	[unknown [Bacillus subtilis]	59	40	603
198	7	5205	4306	gi PID e313073	[hypothetical protein [Bacillus subtilis]	59	38	900
220	5	4362	3958	gi 1PID d101322	[YdhL [Bacillus subtilis]	59	46	405
242	3	1573	2367	gi 1787045	[AE000184] f108: This 108 aa orf is 35 pct identical (15 gaps) to 305 residues of an approx. 296 aa protein PFIC_ECOLI SW: P32675 [Escherichia coli]	59	42	795
247	2	1154	1480	gi 40073	[ORF107 [Bacillus subtilis]	59	39	327

TABLE 2 *S. pneumoniae* - Putative coding regions of novel proteins similar to known proteins

contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
256	1	868	2	[gnl PID d10194	hemolysin [Synechocystis sp.]	59	39	867
258	1	65	820	[g1 2246532	[ORF 73, contains large complex repeat CR 73 [Kaposi's sarcoma-associated herpesvirus]	59	20	756
270	1	386	1126	[gnl PID d10292	[YfnB [Bacillus subtilis]	59	40	741
281	1	552	166	[g1 665062	[putative [Lactococcus lactis]	59	31	387
109	1	3	479	[g1 405879	[yeiH [Escherichia coli]	59	38	477
363	1	2	1894	[g1 915208	[gastric mucin [Sus scrofa]	59	31	1893
387	2	425	84	[g1 160671	[Is antigen precursor [Plasmodium falciparum]	59	44	342
5	6	11233	10465	[gnl PID d101812	[LumQ [Synechocystis sp.]	58	29	759
29	4	2098	3513	[gnl PID d100479	[Na ⁺ - ATPase subunit J [Enterococcus hitrae]	58	39	1416
30	5	4058	3651	[g1 39478	[ATP binding protein of transport ATPases [Bacillus firmus]	58	34	408
33	6	2983	2210	[gnl PID d101164	[unknown [Bacillus subtilis]	58	45	774
36	8	5316	6179	[g1 1538679	[orf [Bacillus subtilis]	58	32	864
43	5	5926	3971	[g1 1788150	[AE000278] protease II [Escherichia coli]	58	37	1956
46	5	3704	5221	[gnl PID e267129	[unknown [Bacillus subtilis]	58	42	1518
48	14	11722	11066	[gnl PID d101771	[chitin biosynthetic bifunctional enzyme [Synechocystis sp.]	58	34	657
52	1	1229	1	[gnl PID d101291	[reductase [Pseudomonas aeruginosa]	58	35	1227
53	2	702	412	[g1 2313357	[AE000545] cytochrome c biogenesis protein (ccdA) [Helicobacter pylori]	58	25	291
58	4	6586	5498	[g1 147329	[transport protein [Escherichia coli]	58	41	1089
69	5	4934	3807	[gnl PID e311492	[unknown [Bacillus subtilis]	58	41	1128
71	27	131357	12277	[g1 2408014	[hypothetical protein [Schizosaccharomyces pombe]	58	33	921
72	4	3586	2882	[g1 18694	[nodulin-21 (AA 1-201) [Glycine max]	58	34	705
74	3	4937	4230	[g1 2293252	[IAF008220] YtmO [Bacillus subtilis]	58	33	708
79	4	4594	3422	[g1 1217989	[ORF3 [Streptococcus pneumoniae]	58	44	1173
82	8	10585	8171	[g1 882711	[exonuclease V alpha-subunit [Escherichia coli]	58	38	2415
86	17	16017	15337	[g1 47642	[5-dehydroquinate hydrolase (3-dehydroquinate) [Salmonella typhi]	58	32	681
97	2	931	560	[g1 153794	[sgg [Streptococcus gordoni]]	58	32	372

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
108	2	358	2724	[g1 537020	[vacB gene product [Escherichia coli]	58	37	2367
111	5	4533	5240	[g1 1592142	[ABC transporter, probable ATP-binding subunit [Methanococcus jannaschii]	58	36	648
120	3	4421	5110	[gml PID d101320	[YggX [Bacillus subtilis]	58	47	690
128	116	13131	12673	[g1 662919	[ORF U [Enterococcus hirae]	58	42	459
132	3	6174	4939	[g1 1800301	[macrolide-efflux determinant [Streptococcus pneumoniae]	58	35	1236
133	1	111	830	[gml PID e269488	[Unknown [Bacillus subtilis]	58	36	780
160	111	8615	9865	[g1 473901	[ORF1 [Lactococcus lactis]	58	39	1251
161	6	6248	6849	[gml PID d101024	[DJ-1 protein [Homo sapiens]	58	32	582
169	1	214	2	[gml PID d100447	[translation elongation factor-3 [Chlorella virus]	58	31	213
187	1	487	2	[g1 475114	[regulatory protein [Pediococcus pentosaceus]]	58	38	486
187	6	4384	4620	[g1 167475	[desiccation-related protein [Craterostigma plantagineum]	58	35	237
190	2	1444	1640	[gml PID e246727	[competence pheromone [Streptococcus gordonii]]	58	38	177
192	2	2012	1344	[gml PID d100556	[rat GCP160 (Rattus rattus)]	58	44	669
206	1	1292	696	[gml PID e202579	[product similar to WbaB [Lactobacillus sakei]]	58	35	597
216	2	2333	555	[gml PID e2425016	[hypothetical protein [Bacillus subtilis]]	58	33	1779
217	5	5250	4321	[g1 466474	[cellobiose phosphotransferase enzyme II' [Bacillus stearothermophilus]]	58	38	930
217	7	5636	5106	[gml PID d102048	[B. subtilis cellobiose phosphotransferase system celB; P46317 (1998) transmembrane [Bacillus subtilis]]	58	44	531
232	1	2	811	[g1 1573777	[cell division ATP-binding protein (ftsE) [Haemophilus influenzae]]	58	39	810
264	1	2	715	[g1 972330	[Nata [Bacillus subtilis]]	58	32	714
280	1	33	767	[g1 1786187	[AB0000111] hypothetical 29.6 kD protein in thrC-talB intergenic region	58	31	715
306	1	845	3	[gml PID e334780	[Yb1 protein [Bacillus subtilis]]	58	47	843
360	3	1556	1092	[sp P46351 Y2GD-	[HYPOTHETICAL 45.4 KD PROTEIN IN THIAMINASE I 5' REGION]	58	32	465
363	5	2160	1867	[g1 160571	[S antigen precursor [Plasmodium falciparum]]	58	51	294
372	1	806	3	[g1 393394	[Tb-291 membrane associated protein [Trypanosoma brucei subgroup]]	58	37	804
382	2	749	519	[pir JC1151 JC11	[hypothetical 20.3K protein (insertion sequence IS1111) - Agrobacterium tumefaciens (strain R022) plasmid Ti]	58	41	231

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
3	9	8409	7471	[gi 1499745	[M. jannaschii predicted coding region MJ0912 [Methanococcus jannaschii]	57	38	939
10	10	7674	7507	[gi 1737169	[homologue to SKP1 [Arabidopsis thaliana]	57	30	168
11	1	2	412	[gnl P101d100139	[ORF [Acetobacter pasteurianus]	57	42	411
31	4	2032	1388	[gi 2293213	[AF008220] YCP1 [Bacillus subtilis]	57	37	645
33	11	6931	6449	[gnl P101d24949	[hypothetical protein [Bacillus subtilis]	57	36	483
45	5	5446	5060	[gi 1592204	[phosphoserine phosphatase [Methanococcus jannaschii]	57	44	387
49	7	6523	7632	[gi 155362	[PTS enzyme-II fructose [Xanthomonas campestris]	57	35	1110
52	6	4520	6850	[gi 1574144	[single-stranded-DNA-specific exonuclease (recJ) [Haemophilus influenzae]	57	35	2331
53	5	2079	1795	[gi 1843180	[replicase-associated polyprotein [oat blue dwarf virus]	57	46	285
63	6	5312	4995	[gi 2182608	[AE000094] YafJ [Rhizobium sp. NGR234]	57	39	318
72	15	13883	11059	[gnl P101d100892	[homologous to SwissProt YIDA_ECOLI hypothetical protein [Bacillus subtilis]	57	40	825
79	2	2561	1815	[gnl P101d100965	[homologue of NADPH-Flavin oxidoreductase Frp of <i>V. harveyi</i> [Bacillus subtilis]	57	44	747
82	9	9596	9763	[gi 1206045	[short region of similarity to glycerophosphoryl diester phosphodiesterases	57	35	168
86	16	15371	14493	[gi 1787983	[AE002641] o288; 92 pct identical (1 gaps) to 222 residues of fragment	57	34	879
93	3	1695	1177	[gi 1500003	[YDB_ECOLI SP: P28244 (223 aa) [Escherichia coli]	57	33	519
96	6	3026	4519	[gi 559882	[mutator mutt protein [Methanococcus jannaschii]	57	43	1494
99	14	17211	16212	[gi 1773349	[threonine synthase [Arabidopsis thaliana]	57	44	1002
112	8	7448	7903	[gi 1591393	[B1B protein [Bacillus subtilis]	57	30	456
113	16	18627	18128	[pir A45605 A45656	[M. jannaschii predicted coding region MJ0578 [Methanococcus jannaschii]	57	22	100
123	2	343	1110	[pir F64149 F641	[hypothetical protein MJ055 - Haemophilus influenzae (strain Rd KW0)	57	38	768
123	4	2108	2884	[gnl P101d102148	[AB001684] sulfate transport system permease protein [Chlorella vulgaris]	57	39	777
127	10	6477	587	[gi 1573082	[nitrogenase C (nifC) [Haemophilus influenzae]	57	35	891
128	13	9251	9790	[gi 153692	[pneumolysin [Streptococcus pneumoniae]	57	38	540
131	4	2139	1363	[gi 42081	[nagI gene product (AA 1-250) [Escherichia coli]	57	16	777

TABLE 2 *S. pneumoniae* - Putative coding regions of novel proteins similar to known proteins

Contig ID	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
136	1	214	1221	bbs 148453	[Spa]endocarditis immunodominant antigen [Streptococcus sobrinus]	57	44	1008
				[263, Peptide, 1566 aa]	[Streptococcus sobrinus]			
140	25	28701	126851	gi 505576	[beta-D-glucoside permease [Bacillus subtilis]	57	38	1851
141	6	6395	7438	gi 995560	unknown [Schizosaccharomyces pombe]	57	41	1044
144	3	3231	2785	gnl P0100139	[ORF] [Acetobacter pasteurianus]	57	42	447
155	4	5454	4564	gi 600411	[glycosyl transferase [Erwinia amylovora]	57	34	891
159	9	4877	5854	gi 290509	[0307 [Escherichia coli]	57	35	978
167	11	9710	9249	gnl P0100139	[ORF] [Acetobacter pasteurianus]	57	42	462
171	6	4023	4436	gi 147402	[mannose permease subunit III-Man [Escherichia coli]	57	29	414
176	4	2170	1076	gnl P0102004	[1AB0014881] ATP-DEPENDENT RNA HELICASE DEAD HOMOLOG. [Bacillus subtilis]	57	39	1095
190	1	145	1455	gi 149420	[export/processing protein [Lactococcus lactis]	57	30	1311
198	1	298	95	gi 522268	[unidentified ORF22 [Bacteriophage blu67]	57	36	204
203	2	3195	2110	gnl P010283915	[orf c01003 [Sulfolobus solfataricus]	57	41	1066
205	1	40	507	gi 1439527	[E11A-man [Lactobacillus curvatus]	57	28	468
214	7	4243	3797	gnl P0102049	[H. influenzae ribosomal protein alanine acetyltransferase, P44305 (1.89) [Bacillus subtilis]	57	48	447
268	3	1767	1276	gi 43979	[L. curvatus small cryptic plasmid gene for rep protein [Lactobacillus curvatus]	57	36	492
351	1	324	34	gnl P010275871	[T03F6_b [Caenorhabditis elegans]	57	31	291
					[S antigen precursor [Plasmodium falciparum]			
386	1	226	2	gi 160671		57	45	225
5	5	10486	8777	gi 405857	[yehU [Escherichia coli]	56	33	170
8	5	3674	3910	gi 467199	[pkSC: LS18_F1_2 [Mycobacterium leprae]	56	39	237
10	3	3442	1874	gnl P0101907	[sodium-coupled permease [Synechocystis sp.]	56	36	1559
21	1	1880	333	gi 2313949	[1AB000593] osmoprotection protein (pr0WX) [Helicobacter pylori]	56	33	1548
22	29	21968	22456	gnl P0102001	[AB001488] PROBABLE ACETYLTRANSFERASE. [Bacillus subtilis]	56	37	489
					[lambda]	56	30	1359
27	1	1361	3	gi 215132	[aa59 (525) [Bacteriophage lambda]	56	41	384
28	9	4667	4278	gi 1592090	[DNA repair protein RAD2 [Methanococcus jannaschii]	56	29	390
33	1	3	386	gnl P0100139	[ORF] [Acetobacter pasteurianus]	56	41	384

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins' similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
36	7	5122	5397	[pir PQ0053 PQ00	hypothetical protein (prod 3' region) - <i>Pseudomonas aeruginosa</i> (strain PAO)	56	28	276
				(fragment)				
40	4	3137	4318	[gi 18060301	macrolide-eflux determinant [<i>Streptococcus pneumoniae</i>]	56	27	1182
40	16	12511	13191	[gnl PID e217602 P1nu	[<i>Lactobacillus plantarum</i>]	56	38	681
48	17	13775	13023	[gi 1143729	transcription activator [<i>Bacillus subtilis</i>]	56	35	753
75	4	1674	2594	[gnl PID d102036 membrane protein	[<i>Bacillus stearothermophilus</i>]	56	25	921
85	3	1802	1459	[gnl PID d100139 ORF	[<i>Acetobacter pasteurianus</i>]	56	41	384
89	7	5815	4940	[gi 853777 product	similar to <i>E. coli</i> PRP2 protein [<i>Bacillus subtilis</i>]	56	42	876
105	2	1360	2718	[gnl PID d101913 hypothetical protein	[<i>Synechocystis</i> sp.]	56	37	1359
112	3	2151	3194	[gi 537201 ORF	_0345 [<i>Escherichia coli</i>]	56	31	1044
113	4	2754	2963	[gnl PID d100340 ORF	[<i>Plum pox</i> virus]	56	28	210
122	3	1203	2054	[gi 1649035 high-affinity periplasmic glutamine binding protein	[<i>Salmonella typhimurium</i>]	56	30	852
124	8	3939	3694	[gnl PID e44893 unknown	[<i>Mycobacterium tuberculosis</i>]	56	27	246
125	4	4403	4107	[gnl PID d100247 human non-muscle myosin heavy chain	[<i>Homo sapiens</i>]	56	32	297
127	111	6608	6005	[gi 12182397 AE00000731 Y4N	[<i>Rhizobium</i> sp. NGR214]	56	35	204
134	5	4769	3849	[gnl PID d101870 hypothetical protein	[<i>Synechocystis</i> sp.]	56	39	921
137	10	6814	7245	[gi 1592011 sulfate permease	[<i>CysA</i>] [<i>Methanococcus jannaschii</i>]	56	34	432
142	18	5019	4582	[pir IA47071 IA470 orf1	immediately 5' of ntrS - <i>Bacillus subtilis</i>	56	29	436
146	8	4676	3660	[gnl PID d101911 hypothetical protein	[<i>Synechocystis</i> sp.]	56	32	1017
148	3	1906	2739	[gnl PID d101059 phosphate transport	system permease protein <i>PstA</i> [<i>Synechocystis</i> sp.]	56	36	834
150	4	4449	2743	[gnl PID e004628 probably site-specific recombinase	of the resolvase family of enzymes	56	27	1707
172	1	2	208	[gi 1787791 AE0002491 f317; This 317 aa	orf is 27 pct identical (16 gaps) to 301 residues of an approx. 320 aa protein YXC_BACSU SW: P39140 [<i>Escherichia coli</i>]	56	34	207
				(f317; This 317 aa	orf is 27 pct identical (16 gaps) to 301 residues of an approx. 320 aa protein YXC_BACSU SW: P39140 [<i>Escherichia coli</i>]	56	40	690
186	7	3732	3367	[gi 1732200 PTS permease	for mannose subunit II ^{Man} [<i>Vibrio furnissii</i>]	56	36	366
187	2	2402	819	[pir IS57941 IS579 virR9	protein - <i>Streptococcus pyogenes</i> (strain CS101, serotype M49)	56	35	1584

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins: similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match gene name	% sim	% ident	length (nt)
				accession			
204	3	2772	2239	gi 6063176 [ORF_0162 (Escherichia coli)]	56	35	534
206	2	3342	1633	gi 559861 [clmA (Plasmid pAD1)]	56	38	1710
219	3	1689	1096	gi 1146197 [putative (Bacillus subtilis)]	56	27	594
230	2	409	1485	pir C60328 C603 [hypothetical protein 2 (str 5' region) - Streptococcus mutans (strain OMZ175 serotype F)]	56	40	1077
233	4	2930	3268	gi 1041785 [Rho-type protein (Plasmid yeast1)]	56	24	339
273	2	1543	2724	gi 143089 [Lep protein (Bacillus subtilis)]	56	32	1182
353	1	1	516	gnl P010325000 [hypothetical protein (Bacillus subtilis)]	56	41	516
359	1	87	641	gi 1786352 [(AE000176) 0877; 100 pct identical to the first 86 residues of the 100 aa hypothetical protein fragment YGBB_ECOLI SW: P54716 (Escherichia coli)]	56	46	555
363	7	4482	4198	gi 1573353 [outer membrane integrity protein (tola) (Haemophilus influenzae)]	56	38	285
376	1	2	508	gnl P010325031 [hypothetical protein (Bacillus subtilis)]	56	33	507
18	1	836	177	gnl P0100872 [a negative regulator of pho regulon (Pseudomonas aeruginosa)]	55	31	660
28	4	1824	1618	gnl P010316518 [STAT protein (Dictyostelium discoideum)]	55	40	207
29	6	4496	5041	gi 1089261 [unknown protein (Anabaena sp.)]	55	31	546
38	16	9695	10702	gi 1580905 [B. subtilis genes rpmI, rpmK, 50kd, gda and gldB (Bacillus subtilis)]	55	31	1008
49	5	5727	6182	gi 1786951 [(AE000176) heat-responsive regulatory protein (Escherichia coli)]	55	29	456
51	4	2381	3241	gnl P0101293 [YbbA (Bacillus subtilis)]	55	42	861
52	9	9640	10866	gi 153016 [ORF_419 protein (Staphylococcus aureus)]	55	23	1227
53	4	1813	1349	gi 896042 [OspF (Borrelia burgdorferi)]	55	30	465
60	5	4794	5756	gi 1499876 [magnesium and cobalt transport protein (Methanococcus jannaschii)]	55	38	963
71	9	14176	15408	gi 11857120 [Glycosyl transferase (Neisseria meningitidis)]	55	41	1233
75	6	3189	4229	gnl P010209890 [NAD alcohol dehydrogenase (Bacillus subtilis)]	55	44	1041
108	10	10488	9820	gnl P010324997 [hypothetical protein (Bacillus subtilis)]	55	36	669
113	12	12273	13037	gnl P010311496 [unknown (Bacillus subtilis)]	55	34	765
113	13	13007	13945	gi 1573423 [1-phosphofructokinase (fruK) (Haemophilus influenzae)]	55	39	939
126	5	6764	5907	gi 1790131 [(AE000446) hypothetical 29.7 kD protein in ibpA-gyrB intergenic region (Escherichia coli)]	55	37	858

TABLE 2 *S. pneumoniae* - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	start (nt)	stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
129	3	2719	902	gnl PID d101425	[<i>Peptidase</i> (<i>Bacillus licheniformis</i>)]	55	35	1818
138	3	2593	1610	gi 142833	[ORF2 (<i>Bacillus subtilis</i>)]	55	37	984
140	6	6916	5633	gnl PID d100964	[homologue of hypothetical protein in a kanamycin synthesis gene cluster of <i>S. pneumoniae</i> hygrosopicus (<i>Bacillus subtilis</i>)]	55	26	1284
147	3	3854	2136	gi 472310	[<i>Hydroxylpeptide dehydrogenase</i> (<i>Clostridium magnum</i>)]	55	39	1719
147	10	110204	8921	gnl PID e73078	[<i>Dihydroorotate</i> (<i>Lactobacillus leichmannii</i>)]	55	38	1284
148	5	3430	4119	gi 1290572	[<i>Peripheral membrane protein U</i> (<i>Escherichia coli</i>)]	55	29	690
148	6	4171	4650	gi 695769	[<i>Transposase</i> (<i>Xanthobacter autotrophicus</i>)]	55	37	480
149	14	112564	11650	gnl PID d101329	[<i>YqjG</i> (<i>Bacillus subtilis</i>)]	55	32	915
156	3	1113	550	gi 12314496	[<i>AE000634</i> conserved hypothetical integral membrane protein (<i>Helicobacter pylori</i>)]	55	34	564
159	10	6625	5897	gi 1290533	[<i>Similar to E. coli</i> ORF adjacent to <i>suc</i> operon; similar to <i>gntr</i> class of regulatory proteins (<i>Escherichia coli</i>)]	55	29	729
164	3	1784	2332	gnl PID e255118	[<i>Hypothetical protein</i> (<i>Bacillus subtilis</i>)]	55	37	549
164	5	2772	3521	gi 140348	[<i>Put</i> . <i>resolvase</i> <i>Tnp I</i> (AA 1 - 284) (<i>Bacillus thuringiensis</i>)]	55	35	750
164	11	7428	7216	gnl PID e249407	[<i>Unknown</i> (<i>Mycobacterium tuberculosis</i>)]	55	38	213
167	5	3860	3345	gi 535052	[<i>Involved in protein secretion</i> (<i>Bacillus subtilis</i>)]	55	28	516
186	5	2880	2563	gi 606080	[<i>ORF0290</i> : <i>Geneplot</i> suggests frameshift linking to 0267, not found in <i>Escherichia coli</i>]	55	35	318
189	8	4311	5396	gnl PID e1833450	[<i>Hypothetical EcS8 protein</i> (<i>Bacillus subtilis</i>)]	55	32	1086
192	5	3270	3079	gi 1196504	[<i>Vitellogenin convertase</i> (<i>Aedes aegypti</i>)]	55	38	192
195	2	2454	1384	gi 1574693	[<i>transfase</i> . <i>Peptidoglycan synthesis</i> (<i>murG</i>) (<i>Haemophilus influenzae</i>)]	55	33	1071
198	4	3013	2471	gnl PID e133074	[<i>Hypothetical Protein</i> (<i>Bacillus subtilis</i>)]	55	29	543
214	1	373	744	gnl PID d101741	[<i>Transposase</i> (<i>Synechocystis</i> sp.)]	55	33	372
219	2	1115	456	gi 288301	[<i>ORF2</i> gene product (<i>Bacillus megaterium</i>)]	55	30	660
263	7	3742	3443	gi 18137	[<i>cgr-4</i> product (<i>Chlamydomonas reinhardtii</i>)]	55	48	300
285	1	2	829	gnl PID d100974	[<i>Unknown</i> (<i>Bacillus subtilis</i>)]	55	40	828
286	1	650	249	gi 396844	[<i>ORF</i> (18 kDa) (<i>Vibrio cholerae</i>)]	55	31	402
297	2	1229	1696	gi 150848	[<i>prtc</i> (<i>Porphyromonas gingivalis</i>)]	55	39	468

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins-similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
309	2	218	982	gi 157491	hypothetical [Haemophilus influenzae]	55	35	765
328	2	646	224	gi 57150	prohibitin [Saccharomyces cerevisiae]	55	27	423
330	1	1340	474	gi 396197	soxS [Escherichia coli]	55	29	867
364	3	2538	1546	gi 1393394	Tb-291 membrane associated protein [Trypanosoma brucei subgroup]	55	36	993
368	3	941	105	gi 1160671	IS antigen precursor [Plasmodium falciparum]	55	40	837
3	5	4604	3624	gi 12293176	[AF008220] signal transduction protein kinase [Bacillus subtilis]	54	26	981
9	11	7746	7246	gi 11466245	putative [Bacillus subtilis]	54	38	501
38	124	16213	17937	gi 1400429	putative transcriptional regulator [Bacillus stearothermophilus]	54	27	1725
40	8	5076	4682	gi 39969	methionyl-tRNA synthetase [Bacillus stearothermophilus]	54	35	195
43	4	3980	2267	gnl PID el48611	ABC transporter [Lactobacillus helveticus]	54	25	1614
52	10	10844	12103	gi 11702962	FemA [Staphylococcus simulans]	54	29	1260
57	1	3	512	gi 558177	endo-1,4-beta-D-xylosidase [Cellulomonas fimi]	54	36	510
58	3	4749	4246	gnl PID d101237	hypothetical [Bacillus subtilis]	54	29	504
71	7	10884	11703	gi 1510255	Orf3 [Escherichia coli]	54	31	1020
71	120	127546	127337	gi 1202543	serotonin receptor [Rattus norvegicus]	54	31	192
72	2	804	1098	gi 148613	srnB gene product [Plasmid F]	54	37	255
72	7	7438	6695	gi 1196466	recombinase [Moxxella bovis]	54	38	744
74	10	14043	112465	gi 1200312	[ORF 3 gene product [Bradyrhizobium japonicum]	54	32	579
74	12	16683	15995	gi 2317798	[maturase-related protein [Pseudomonas alcaligenes]	54	10	489
86	3	2877	2155	gi 46988	[orf9.6 possibly encodes the O unit polymerase [Salmonella enterica]	54	34	723
89	5	4333	3921	gi 147211	[phoD protein [Escherichia coli]	54	41	513
90	1	3	464	gi 2317798	[maturase-related protein [Pseudomonas alcaligenes]	54	30	462
96	10	8058	8510	gnl PID d102015	[AB001488] SIMILAR TO SALMONELLA TYPHIMURUM SLT7 GENE REQUIRED FOR SURVIVAL IN MACROPHAGE. [Bacillus subtilis]	54	32	453
97	6	4662	3604	gi 1591394	[transketolase] [Methanococcus jannaschii]	54	30	1059
106	11	10406	12010	gi 606286	[ORF_0637] [Escherichia coli]	54	32	1605
147	8	8663	7404	gnl PID d101615	[ORF_ID:031947; similar to SwissProt Accession Number P37340] [Escherichia coli]	54	35	1260

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
171	4	2477	3223	[gi 1439328	[E. coli-mn [Lactobacillus curvatus]	54	36	747
174	2	2068	1787	[gnl PID1d00518	[motor protein [Homo sapiens]	54	35	282
188	1	526	1188	[gnl PID1e250352	[unknown [Mycobacterium tuberculosis]	54	31	663
198	5	3582	2884	[gnl PID0e313074	[hypothetical protein [Bacillus subtilis]	54	33	699
207	1	1	1641	[gnl PID1d01813	[hypothetical protein [Synochocystis sp.]	54	24	1641
210	1	2	655	[gi 2293206	[AF008220] Ymp [Bacillus subtilis]	54	29	654
225	2	966	2357	[gnl PID0e310194	[R11H6.1 [Caenorhabditis elegans]	54	39	1392
241	1	1681	347	[gnl PID1d01813	[hypothetical protein [Synochocystis sp.]	54	26	1335
263	2	907	1395	[gnl PID0f01886	[transposase [Synochocystis sp.]	54	30	489
263	6	3450	2977	[gi 160671	[S antigen precursor [Plasmodium falciparum]	54	47	474
277	3	2517	1363	[gi 11195926	[unknown protein [Streptococcus mutans]	54	30	1155
307	1	828	4	[gi 2293198	[AF008220] Ycp [Bacillus subtilis]	54	28	825
325	1	19	768	[gi 2182507	[AE0000083] Y41H [Rhizobium sp. NGR234]	54	37	750
332	2	898	590	[gi 1591815	[ADP-ribosylglycohydrolase (draG) [Methanococcus jannaschii]	54	32	309
385	4	240	479	[gi 530888	amino acid feature: N-glycosylation sites, aa 41 .. 43, 46 .. 48, 51 .. 53, 72 .. 74, 107 .. 109, 128 .. 130, 132 .. 134, 158 .. 160, 163 .. 165; amino acid feature: Rod protein domain, aa 169 .. 340; amino acid feature: globular protein domain	54	49	240
7	25	19702	19493	[gnl PID1e255111	[hypothetical protein [Bacillus subtilis]	53	32	210
23	3	2457	2033	[gnl PID1d02015	[AB001488] SIMILAR TO SALMONELLA TYPHIMURIUM SLTIV GENE REQUIRED FOR SURVIVAL IN MACROPHAGE. [Bacillus subtilis]	53	25	465
29	111	9042	10121	[gi 143331	[alkaline phosphatase regulatory protein [Bacillus subtilis]	53	31	1080
33	3	1479	1009	[pir S1065 S106	[hypothetical protein X - Pyrococcus woesei (fragment)	53	33	471
36	6	4383	5134	[gnl PID1e3116029	[unknown [Mycobacterium tuberculosis]	53	30	552
38	14	8821	8898	[gi 580904	[homologous to E. coli rnpA [Bacillus subtilis]	53	30	378
52	7	7007	8686	[gi 1377831	[unknown [Bacillus subtilis]	53	29	1680
54	17	11755	119564	[gi 666069	[orf2 gene product [Lactobacillus leichmannii]	53	36	2010
56	1	1	681	[gi 1592266	[restriction modification system S subunit [Methanococcus jannaschii]	53	32	681

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins-similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
57	10	9431	8487	gi 1788343	(NE000310) f351: Residues 1-121 are 100 pct identical to YQJL_ECOLI SW: P33944 (122 aa) and aa 152-351 are 100 pct identical to YQJK_ECOLI SW: P33943 (Escherichia coli)	53	31	945
61	1	429	4	gi PID e236467	[caenorhabditis elegans]	53	33	426
71	1	5772	4	gi 1393394	[Tb-291] membrane associated protein [Trypanosoma brucei subgroup]	53	33	5769
72	3	894	2840	gi 2293178	[AF008220] Ytsd (Bacillus subtilis)	53	27	1947
73	14	9793	9212	gi 1777556	[putative cobalamin synthesis protein [Escherichia coli]]	53	32	582
88	7	5217	4342	gi 2098719	[putative fimbrial-associated protein [Actinomyces naeslundii]]	53	38	876
93	5	2395	1688	gi 563366	[gluconate oxidoreductase [Gluconobacter oxydans]]	53	33	708
96	9	6632	7762	gi 517204	[ORF1, putative 42 kDa protein [Streptococcus pyogenes]]	53	42	1131
108	8	7629	8600	gi 1449581	[maturation protein [Lactobacillus paracasei]]	53	32	972
128	9	6412	6972	gi PID e317237	[unknown [Mycobacterium tuberculosis]]	53	36	561
128	112	8429	9253	gi 311070	[penetraxin fusion protein [Xenopus laevis]]	53	31	825
148	1	3	950	lpir A1607 A1616	[probable hemolysin precursor - Streptococcus agalactiae (strain 74-360)]	53	36	948
163	2	2162	3022	gi 1775150	[nocurnin [Xenopus laevis]]	53	30	861
171	3	2304	2624	gi 17732200	[PTS permease for mannose subunit IIIPMan [Vibrio furnissii]]	53	32	321
182	5	3785	3051	gi PID d100572	[unknown [Bacillus subtilis]]	53	35	735
209	3	2948	1935	gi 1778505	[ferric enterobactin transport protein [Escherichia coli]]	53	28	1014
218	5	3884	2406	gi 40162	[murE gene product [Bacillus subtilis]]	53	34	1479
250	3	473	790	gi PID e334776	[Ybh protein [Bacillus subtilis]]	53	30	318
275	1	1	1611	gi PID d101314	[Yqaw [Bacillus subtilis]]	53	35	1611
332	1	544	2	gi 409286	[bmrU [Bacillus subtilis]]	53	31	543
2	2	2543	3445	gi PID e233879	[hypothetical protein [Bacillus subtilis]]	52	39	903
3	122	22402	23376	gi 35969	[lacF gene product [Agrobacterium radiobacter]]	52	36	975
5	3	8094	2356	gi PID e324915	[IgM protease [Streptococcus sanguis]]	52	32	579
1	22	13961	120212	gi 152901	[ORF 3 [Spirochaeta aurantia]]	52	35	232
22	31	23140	124666	gi 269262	[cone ORF3 [Bacillus subtilis]]	52	32	1527
27	6	5397	4801	gi 39573	[P20 (AA 1-178) [Bacillus licheniformis]]	52	35	597

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF	Start	Stop	match	match gene name	% sim	% ident	length (nt)
ID	ID	(nt)	(nt)	accession				
35	10	8604	7357	gi 508241	putative O-antigen transporter [Escherichia coli]	52	27	1248
45	4	4801	3662	gnl PID d102243	(AB005541) homologs are found in E. coli and H. influenzae; see SWISS_PROT ACCB: P42100 [Bacillus subtilis]	52	36	1140
48	18	11485	13726	gnl PID e205174	orf2 [Lactobacillus helveticus]	52	25	660
49	4	5321	5755	gi 2117740	(AF013987) nitrogen regulatory TIA protein [Vibrio cholerae]	52	19	435
54	4	2773	4668	gi 1500472	predicted coding region M1577 [Methanococcus jannaschii]	52	36	1896
54	6	5250	4969	gi 21182453	[M. jannaschii] predicted coding region M1577 [Methanococcus jannaschii]	52	40	282
66	6	8400	6955	gi 431140	[Rhizobium sp. NGR234]	52	30	1446
71	126	130659	131312	gnl PID e314993	unknown [Mycobacterium tuberculosis]	52	27	639
75	2	1673	1035	gnl PID d102271	[AEG00079] Y410 [Streptomyces sp.]	52	32	1455
81	3	1439	2893	gnl PID e3111458	rihamnulose kinase [Bacillus subtilis]	52	37	795
81	8	4987	5781	gi 147403	mannose permease subunit II-P-Man [Escherichia coli]	52	37	1167
83	21	20687	21853	gi 1433365	phosphotriboyl amideimidecarboxylase II (PUR-K; ttg start codon) [Bacillus subtilis]	52	26	1194
86	6	5785	4592	gi 1276879	[Streptococcus thermophilus]	52	26	1530
86	120	119390	17861	gi 1454844	EPSP [Streptococcus thermophilus]	52	33	882
96	13	110540	9659	gi 1288299	ORF 3 [Ishikistosoma mansonii]	52	27	2025
111	1	2	2026	gi 148309	lycylsain B transport protein [Enterococcus faecalis]	52	33	711
112	2	1457	2167	gi 471234	[orf1] [Haemophilus influenzae]	52	33	567
118	3	2931	2365	lbs 151233	[Mip=24 kDa macrophage infectivity potentiator protein [Legionella pneumophila]]	52	16	306
122	1	9	5646	5951	[gi 8214] [Drosophila melanogaster]	52	52	216
122	11	6159	6374	gi 434025	[dihydrofolate acetyltransferase [Pseudomonas carbinolicus]]	52	43	1434
134	6	4880	6313	gi 153733	[M protein trans-acting positive regulator [Streptococcus pyogenes]]	52	35	1479
135	3	1238	2116	gnl PID e245024	[unknown [Mycobacterium tuberculosis]]	52	32	639
141	3	1681	2319	gnl PID d100573	[unknown [Bacillus subtilis]]	52	36	2463
161	4	2562	5024	gi 1146243	22.4% identity with Escherichia coli DNA-damage inducible protein ...; putative [Bacillus subtilis]	52	30	786
173	2	968	183	gi 1215693	[putative orf; G99-orf434 [Mycoplasma pneumoniae]]	52	1	+

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
198	6	4400	3567	[gnl PID e311010	[hypothetical protein [Bacillus subtilis]	52	26	834
				[hypothetical protein [Bacillus subtilis]		52	38	264
210	112	8844	9107	[gi 487647	[DNA gyrase subunit B [Mycoplasma genitalium]	52	36	168
				[DNA gyrase subunit B [Mycoplasma genitalium]		52	36	168
214	110	5264	5431	[gi 550697	[envelope protein [Human immunodeficiency virus type 1]	52	34	870
				[envelope protein [Human immunodeficiency virus type 1]		52	34	870
225	1	15	884	[gi 11552773	[hypothetical [Escherichia coli]	52	28	324
				[hypothetical [Escherichia coli]		52	28	324
230	1	19	362	[gnl PID d100582	[unknown [Bacillus subtilis]	52	29	870
				[unknown [Bacillus subtilis]		52	29	870
287	1	871	2	[gnl PID e335028	[protease/peptidase [Mycobacterium leprae]	52	32	1302
				[protease/peptidase [Mycobacterium leprae]		52	32	1302
163	2	1305	4	[gnl 393394	[Tb-291 membrane associated protein [Tyrpanosoma brucei subgroup]	51	30	876
				[Tb-291 membrane associated protein [Tyrpanosoma brucei subgroup]		51	30	876
23	2	2048	1173	[gnl PID e354943	[unknown [Mycobacterium tuberculosis]	51	31	780
				[unknown [Mycobacterium tuberculosis]		51	31	780
29	3	742	1521	[gi 1929900	[5'-methylthioadenosine phosphorylase [Sulfolobus solfataricus]	51	32	1188
				[5'-methylthioadenosine phosphorylase [Sulfolobus solfataricus]		51	32	1188
45	1	410	1597	[gi 1877429	[Integrase [Streptococcus pyogenes phage T12]	51	33	282
				[Integrase [Streptococcus pyogenes phage T12]		51	33	282
48	126	119227	118946	[gnl PID e2314455	[transcriptional regulator [tenA] [Helicobacter pylori]	51	31	261
				[transcriptional regulator [tenA] [Helicobacter pylori]		51	31	261
73	5	4276	4016	[gi 1474177	[alpha-D-1,4-glucosidase [Staphylococcus xylosus]	51	31	3123
				[alpha-D-1,4-glucosidase [Staphylococcus xylosus]		51	31	3123
81	111	8935	12057	[gi 1111070	[penetrin fusion protein [Xenopus laevis]	51	33	792
				[penetrin fusion protein [Xenopus laevis]		51	33	792
83	5	1195	1986	[gnl PID d101316	[Yqfr [Bacillus subtilis]	51	28	1008
				[Yqfr [Bacillus subtilis]		51	28	1008
98	10	17531	18538	[gi 41500	[orf 3 (AA 1-1521), 38 kd (put. ftsX) [Escherichia coli]	51	27	1266
				[orf 3 (AA 1-1521), 38 kd (put. ftsX) [Escherichia coli]		51	27	1266
113	6	3908	5173	[gi 466882	[ppal: B1496_C2_189 [Mycobacterium leprae]	51	32	270
				[ppal: B1496_C2_189 [Mycobacterium leprae]		51	32	270
124	1	326	57	[gi 2191168	[AF001270] contains similarity to myosin heavy chain [Azobidopsis thaliana]	51	30	471
				[AF001270] contains similarity to myosin heavy chain [Azobidopsis thaliana]		51	30	471
129	10	7286	6816	[gi 1046241	[orf14 [Bacteriophage HP1]	51	26	981
				[orf14 [Bacteriophage HP1]		51	26	981
143	3	4963	3983	[gi 1135935	[probable copper-transporting atpase [Escherichia coli]	51	36	1134
				[probable copper-transporting atpase [Escherichia coli]		51	36	1134
148	15	111359	10226	[gi 12291256	[AF008220] putative hippurate hydrolase [Bacillus subtilis]	51	21	1311
				[AF008220] putative hippurate hydrolase [Bacillus subtilis]		51	21	1311
149	8	6003	7313	[gi 1633572	[Herpesvirus saimiri ORF73 homolog [Kaposi's sarcoma-associated herpesvirus]	51	34	543
				[Herpesvirus saimiri ORF73 homolog [Kaposi's sarcoma-associated herpesvirus]		51	34	543
151	9	112092	11550	[gnl PID e281580	[hypothetical 40-7 kd protein [Bacillus subtilis]	51	36	654
				[hypothetical 40-7 kd protein [Bacillus subtilis]		51	36	654
159	6	2555	3208	[gi 146944	[CDP-N-acetylneuramnic acid synthetase [Escherichia coli]	51	28	1794
				[CDP-N-acetylneuramnic acid synthetase [Escherichia coli]		51	28	1794
174	1	1797	4	[gi 1773166	[probable copper-transporting atpase [Escherichia coli]	51	18	459
				[probable copper-transporting atpase [Escherichia coli]		51	18	459
265	4	2231	1773	[gnl PID e256400	[anti-P-falciparum antigenic polypeptide [Salinaria sciureus]	51	33	669
				[anti-P-falciparum antigenic polypeptide [Salinaria sciureus]		51	33	669
277	2	643	1311	[pir S32915 S329	[pild protein - Neisseria gonorrhoeae	51	33	+
				[pild protein - Neisseria gonorrhoeae]		51	33	+

TABLE 2 S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
350	1	890	3	gi 290569	gi 307 [Escherichia coli]	51	30	888
363	4	1228	4485	gi 1707247	partial CDS [Caenorhabditis elegans]	51	23	3258
367	1	1701	4	gi 1393194	1b-291 membrane associated protein [Trypanosoma brucei subgroup]	51	32	1698
15	5	5174	4497	gi PID e58151	F3 [Bacillus subtilis]	50	38	678
16	4	2220	2582	gnl PID e325010	hypothetical protein [Bacillus subtilis]	50	29	163
19	5	2591	4159	gi 1552733	similar to voltage-gated chloride channel protein [Escherichia coli]	50	30	1569
25	4	2701	1997	gi 887849	ORF_f219 [Escherichia coli]	50	27	705
35	1	211	417	gi PID e235697	unknown [Saccharomyces cerevisiae]	50	33	207
39	4	3416	5152	gi PID d100974	unknown [Bacillus subtilis]	50	27	1737
51	7	4000	5181	gi 1592027	carbamoyl-phosphate synthase, pyrimidine-specific, large subunit [Methanococcus jannaschii]	50	27	1182
51	9	7179	8303	gi 1591847	type I restriction-modification enzyme, S subunit [Methanococcus jannaschii]	50	28	1125
52	8	8740	9534	gi 144297	acyl esterase (XynC) [Candidocellum saccharolyticum]	50	34	795
52	16	16591	15770	gi 2108229	basic surface protein [Lactobacillus fermentum]	50	34	822
57	7	6031	6336	gi 2275264	60S ribosomal protein L7B [Schiizosaccharomyces pombe]	50	40	106
71	23	29348	28383	gi PID d101328	YQPA [Bacillus subtilis]	50	30	966
86	12	11155	110769	gnl PID e3249564	hypothetical protein [Bacillus subtilis]	50	24	387
93	2	1205	330	gi 1066016	similar to Escherichia coli pyruvate, water dikinase, Swiss-Prot Accession Number P21538 [Pyrococcus furiosus]	50	24	876
96	5	1673	2959	gnl PID e322433	gamma-glutamylcysteine synthetase [Brassica juncea]	50	29	1287
98	2	218	1171	gi 151110	[Pseudomonas aeruginosa]	50	30	954
103	4	3303	2785	gi 154330	o-antigen ligase [Salmonella typhimurium]	50	31	519
115	5	6480	5980	gi 895747	putative cel operon regulator [Bacillus subtilis]	50	26	501
129	11	7559	7305	gi 121645	leucine-, isoleucine-, and valine-binding protein [Bacillus subtilis]	50	32	255
129	13	8192	7965	gi 1152271	skeletal muscle ryanodine receptor [Homo sapiens]	50	30	220
151	5	7624	6819	gi 40348	[put. resolvase Rmp I (AA 1 - 284) [Bacillus thuringiensis]	50	35	816
153	1	1	597	gnl PID d102015	1319 kDa protein [Rhizobium meliloti] SIMILAR TO NITROREDUCTASE. [Bacillus subtilis]	50	29	597

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (Int)	Stop (Int)	match accession	match gene name	% sim	% ident	length (Int)
155	5	5986	5432	gi 1276880	[EpSG (Streptococcus thermophilus)]	50	28	555
160	9	7390	6323	gi 1786983	[AE000179] o131: 92 pct identical to the 333 aa hypothetical protein (7 gaps) to 167 residues of the YBHE_ECOLI SW: PS2697; 26 pct identical (7 gaps) to 373 aa protein MLE_TRICU SW: P46057; SW: PS2697 [Escherichia coli]	50	30	1068
163	6	7396	8091	gi 1201313	[YQEN (Bacillus subtilis)]	50	22	696
167	6	5232	3940	gi 413926	lipA-2r gene product [Bacillus subtilis]	50	27	1293
169	2	807	130	gi 100680	[ORF_o290; Geneplot suggests frameshift linking to o267, not found]	50	35	678
171	5	3168	4025	gi 606080	[Endolysin (Bacteriophage Bastille)]	50	27	858
210	111	8151	8414	gi 330038	[HRV 2 polyprotein [Human rhinovirus]	50	25	264
364	1	1538	135	gi 139336	[rb-292 membrane associated protein [Trypanosoma brucei subgroup]	50	31	1404
10	17	5911	5090	gi 144859	[ORF_B [Clostridium perfringens]	49	24	822
26	5	10754	9768	gi 142440	[ATP-dependent nuclease [Bacillus subtilis]]	49	31	987
66	7	9777	8398	gi 414170	[trkA gene product [Methanocarcina mazei]]	49	26	1380
77	6	5364	4648	gi 10285322	[RecX protein [Mycobacterium smegmatis]]	49	28	717
82	113	12689	13249	gi 10255091	[hypothetical protein [Bacillus subtilis]]	49	20	561
93	9	4866	4531	gi 40067	[X gene product [Bacillus sphaericus]]	49	26	316
112	5	4019	4948	gi 1574380	[luc-1 operon protein [lucB] [Haemophilus influenzae]]	49	27	930
129	7	6058	4949	gi 1010 e267587	[Unknown [Bacillus subtilis]]	49	35	1110
135	5	3875	4438	gi 39573	[P20 (AA 1-178) [Bacillus licheniformis]]	49	25	564
154	2	1423	1953	gi 1010 e10102	[regulatory components of sensory transduction system [Synechocystis sp.]]	49	29	531
156	5	2878	1637	gi 1010 e101732	[hypothetical protein [Synechocystis sp.]]	49	25	1242
173	5	3500	2940	gi 490324	[LORF_X gene product [unidentified]]	49	30	561
182	1	1057	2	gi 331002	[first methionine codon in the ECLF1 ORF [Salmarinine herpesvirus 2]]	49	25	1056
192	6	5352	3667	gi 2394472	[AF024499) contains similarity to homeobox domains [Caenorhabditis elegans]]	49	23	1686
253	4	1129	1150	gi 4931116	[SIR4 protein [Saccharomyces cerevisiae]]	49	23	222
277	1	600	136	gi 396844	[ORF (18 kDa) [Vibrio cholerae]]	49	32	465
327	3	1435	887	gi 733524	[phosphatidylinositol-4,5-diphosphate 3-kinase (Dictyostelium discoideum)]	49	24	549

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
365	3	1436	132	[gi 333394	[rb-291 membrane associated protein [Trypanosoma brucei subgroup]	49	31	1305
33	7	4461	3277	[gi 145644	[codes for a protein of unknown function [Escherichia coli]	48	26	1185
40	2	652	1776	[gnl PID e290649	[ornithine decarboxylase [Nicotiana tabacum]	48	29	1125
67	4	1377	2384	[gi 1772652	[2-keto-3-deoxygluconate kinase [Haloflexax alicantae]	48	30	1008
74	2	4269	3871	[gi 2182678	[AS000101] Y4W [Rhizobium sp. NGR234]	48	27	399
81	2	1326	541	[gi 153672	[lactose repressor [Streptococcus mutans]	48	33	786
81	4	2981	3646	[gi 146042	[fuculose-1-phosphate aldolase (fucA) [Escherichia coli]	48	30	666
97	1	602	51	[gi 153794	[rgg [Streptococcus gordonii]	48	29	552
110	1	1	3132	[gi 1381114	[prtb gene product [Lactobacillus delbrueckii]	48	23	3132
131	5	2914	2147	[gnl PID e183811	[Acyl-ACP thioesterase [Brassica napus]	48	27	768
133	4	3494	2628	[gnl PID e261988	[putative ORF [Bacillus subtilis]	48	27	857
139	6	4231	4599	[gi 1049388	[2K47.1 gene product [Caenorhabditis elegans]	48	23	369
139	8	5036	5665	[gi 1022725	[unknown [Staphylococcus haemolyticus]	48	29	630
140	12	11936	11007	[gnl PID d102049	[H. influenzae, ribosomal protein alanine acetyltransferase; P4305 (189)	48	27	930
146	9	5670	4654	[gi 1591731	[malvalonate kinase [Methanococcus jannaschii]	48	24	1017
161	3	1280	2374	[gnl PID d101578	[Collagenase precursor (EC 3.4.-.-) [Escherichia coli]	48	24	1095
172	11	10281	11048	[gnl PID d101132	[hypothetical protein [Synchocystis sp.]	48	27	468
182	4	2330	2586	[gi 40067	[X gene product [Bacillus sphaericus]	48	37	345
210	15	10786	11196	[sp P11940 LE129-	[LATE EMBRYOGENESIS ABUNDANT PROTEIN D-29 (LEA D-29).	48	30	411
214	12	6231	6482	[gi 40389	[non-toxic components [Clostridium botulinum]	48	26	252
221	1	704	3	[gi 1573364	[H. influenzae predicted coding region H10392 (Haemophilus influenzae)	48	27	702
227	2	647	3928	[gi 1673691	[Mycoplasma pneumoniae, C09_orf13 Protein [Mycoplasma pneumoniae]	48	30	3282
253	2	480	758	[gnl PID e236697	[unknown [Saccharomyces cerevisiae]	48	31	279
363	3	1874	1122	[gi 18137	[cgrc-4 product [Chlamydomonas reinhardtii]	48	40	753
389	1	505	2	[gi 18137	[cgrc-4 product [Chlamydomonas reinhardtii]	48	38	504
3	121	120879	12258	[gnl PID e264778	[putative maltose-binding protein [Streptomyces coelicolor]	47	33	1380

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	start (nt)	stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
6	4	4089	4658	[gi 39573]	[P20 (AA 1-178) [Bacillus licheniformis]	47	23	570
15	13	1736	1760	[ORF ID:00572]	[Unknown [Bacillus subtilis]]	47	25	1977
35	115	14516	13263	[gi 1773351]	[Cap5 (Staphylococcus aureus)]	47	20	1254
51	6	3547	4002	[pir A37024 A370]	[32K antigen precursor - Mycobacterium tuberculosis]	47	38	456
55	8	10154	9273	[gi 39848]	[U3 [Bacillus subtilis]]	47	26	882
92	4	1753	3276	[grn PID0le280611]	[PPC (Streptococcus pneumoniae)]	47	35	1524
127	9	5589	5386	[gi 1786458]	(AE000134) f120: this 120 aa orf is 76 pct identical (0 gaps) to 42 residues of an approx. 48 aa protein Y127_HAEIN SW: P43949 [Escherichia coli]	47	32	204
130	2	1232	1759	[grn PID0le265555]	[Unknown [Mycobacterium tuberculosis]]	47	23	528
140	4	4951	3542	[grn PID d100964]	homologue of hypothetical protein in a rhamycin synthesis gene cluster of Streptomyces hygroscopicus [Bacillus subtilis]	47	24	1410
151	4	6814	6200	[gi 1522674]	[M. jannaschii predicted coding region MJECI41 [Methanococcus jannaschii]]	47	27	615
157	3	803	1174	[grn PID d101320]	[Yq92 [Bacillus subtilis]]	47	25	372
178	5	3267	2155	[gi 2367190]	(AE000390) o334: sequence change joins ORFs y9jR & y9jS from earlier version (YGR_ECOLI SW: P42599 and YGTS_ECOLI SW: P42600) [Escherichia coli]	47	30	1113
273	1	2	1549	[grn PID e254973]	[autolysin sensor kinase [Bacillus subtilis]]	47	32	1548
300	2	880	644	[gi 1635755]	[zinc finger protein Png-1 [Mus musculus]]	47	22	237
54	14	14182	12638	[pir sa360 sa36]	[rofa protein - Streptococcus pyogenes]	46	24	1545
88	1	2	1018	[grn PID e223891]	[xylose repressor (Ansarcocellum thermophilum)]	46	20	1017
96	7	4553	5860	[grn PID d101652]	(ORF ID:034785; similar to [Swissprot Accession Number P45272] [Escherichia coli])	46	23	1308
112	1	1127	3	[gi 2209215]	(AF004125) putative oligosaccharide repeat unit transporter [Streptococcus pneumoniae]	46	24	1125
122	13	7308	7982	[gi 1054776]	[hr46 gene product (Homo sapiens)]	46	34	675
127	14	9198	8125	[gi 1469286]	[afua gene product (Actinobacillus pleuropneumoniae)]	46	28	1074
132	4	7093	6197	[gi 53794]	[rgg (Streptococcus gordonii)]	46	26	897
140	8	8220	7723	[gi 1235795]	[pullulanase (Thermanaerobacterium thermosulfurigenes)]	46	21	498
140	9	9205	8315	[gi 40878]	[leucine rich protein (Streptococcus equisimilis)]	46	27	891

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)	match accession	match gene name	% sim	% ident	length (nt)
162	1	1	1125	gi 1143209	[ORF7: Method: conceptual translation supplied by author [Shigella sonnei]	46	25	1125
199	1	1	585	gi 1947171	[AP000299] No definition line found [Caenorhabditis elegans]	46	28	585
223	3	1971	1477	sp P02563 WYSS_	[WYSS: MYOSIN HEAVY CHAIN, SKELETAL MUSCLE (FRAGMENTS)]	46	27	495
232	2	760	1608	gi 1016112	[ycf38 gene product [Cyanophora paradoxa]]	46	28	849
292	1	687	220	gi 1673744	[AE000011] Mycoplasma pneumoniae, cytidine deaminase; similar to GenBank Accession Number C5312, from <i>M. pneumoniae</i>	46	29	468
30	8	5843	6472	gi 1788049	[AE000270] 0235; This 235 aa orf is 29 pct identical (10 gops) to 198 residues of an approx. 216 aa protein YTB_BACSU SW: P06568 [Escherichia coli]	45	24	630
48	6	3461	3868	gi 722339	[Unknown [Acetobacter xylinum]]	45	29	408
60	1	307	2	gi 1690909	[coded for by <i>C. elegans</i> cDNA yk41h4.3; coded for by <i>C. elegans</i> cDNA yk14g10.5; coded for by <i>C. elegans</i> cDNA yk152g5.5; coded for by <i>C. elegans</i> cDNA yk39a10.5; coded for by <i>C. elegans</i> cDNA yk41h4.5; coded for by <i>C. elegans</i> cDNA cm20g10; coded for by <i>C. elegans</i> cDNA cm20g10]	45	36	306
72	16	14371	14874	gi 1321930	[NADH dehydrogenase (ubiquinone) (Artemia franciscana)]	45	25	504
99	7	9158	7941	gi 152192	[mutation causes a succinoglucan-minus phenotype: ExoQ is a transmembrane protein; third gene of the exoFQ operon.; putative [Rhizobium meliloti]]	45	28	1218
127	12	7046	6606	bhs 153689	[HtrB-iron utilization protein [Haemophilus influenzae, type b, DL42, NTHI strain 106. Peptide, 506 aa]; Haemophilus influenzae]	45	24	441
137	5	1561	2619	gi 1472921	[v-type Na-ATPase (Enterococcus hirae)]	45	33	1059
209	1	774	364	gi 1304141	[restriction endonuclease beta subunit [Bacillus coagulans]]	45	28	411
314	1	604	2	gi 1400457	[Latex allergen [Havea brasiliensis]]	45	31	603
20	116	19782	20388	gi 433942	[ORF [Lactococcus lactis]]	44	26	507
87	8	7030	6452	gi 537207	[ORF f277 [Escherichia coli]]	44	26	579
166	5	4909	4037	gi P08002	[membrane transport protein [Bacillus subtilis]]	44	25	873
247	1	818	75	gi P00718	[ORF1 [Bacillus sp.]]	44	20	744
32	3	1885	3876	gi 2351768	[PsPa [Streptococcus pneumoniae]]	43	24	1992
36	17	15467	18256	gi 1045739	[M. genitalium predicted coding region W0054 [Mycoplasma genitalium]]	43	26	2790
54	15	14656	17343	gi 1520341	[penicillin-binding proteins 1A and 1B [Bacillus subtilis]]	43	27	2688
67	1	696	1352	gi 536934	[yjCA gene product [Escherichia coli]]	43	29	657
139	2	2416	338	gi 39600	[similar to eukaryotic Na ⁺ /H ⁺ exchangers [Escherichia coli]]	43	24	2079

TABLE 2
S. pneumoniae - Putative coding regions of novel proteins similar to known proteins

Contig	ORF ID	Start ID	Stop ID	match accession	match gene name	% sim	% ident	length (nt)
298	1	3	809	gi 1413972	lipo-08r gene product [Bacillus subtilis]	43	24	807
387	1	47	427	gi 231552	(AF016669) No definition line found [Caenorhabditis elegans]	43	30	381
165	4	4221	3127	gi 12182399	(AE000073) Y4FP [Rhizobium sp. NGR234]	41	25	1095
340	1	582	70	gnl PID e218681	CDP-diacylglycerol synthetase [Arabidopsis thaliana]	41	20	513
363	6	4205	1914	gi 1256742	R27-2 protein [Trypanosoma cruzi]	41	27	2292
368	2	2	943	gi 21703	LWY glutinin (AA 1-356) [Triticum aestivum]	41	34	942
155	3	4489	2861	gi 42023	member of ATP-dependent transport family, very similar to mdr proteins and hemolysin B, export protein [Escherichia coli]	40	18	1629
365	2	95	1438	gi 1633572	Herpesvirus saimiri ORF73 homolog [Kaposi's sarcoma-associated herpes-like virus]	40	21	1344
1	3	2979	3860	gnl PID d101908	hypothetical protein [Synechocystis sp.]	39	26	882
1	5	3814	4647	gnl PID d101961	hypothetical protein [Synechocystis sp.]	39	19	634
26	6	14035	10724	gi 114239	ATP-dependent nuclease [Bacillus subtilis]	38	20	3112
47	1	3	4916	gi 632549	INF-180 [Petcomyzon marinus]	36	23	4914

TABLE 3
S. pneumoniae - Putative coding regions of novel proteins not similar to known proteins

Contig	ORF	Start (nt)	Stop (nt)
ID	ID	(nt)	(nt)
1	4	3428	3009
1	6	4611	4964
3	2	818	994
3	3	1182	1574
3	7	5382	6497
3	25	25046	25396
3	26	125625	126317
6	2	1519	1689
6	14	12875	12616
6	15	13215	12841
6	18	15977	15390
7	12	9935	9419
7	13	10161	9910
8	6	3915	4280
9	9	6024	5704
10	8	6909	6298
10	9	7136	6888
10	11	7968	7672
12	1	1140	4
12	3	1779	1456
14	2	1913	1434
16	1	1	243
16	5	5675	3087
17	1	324	34
17	3	1451	1050
17	9	4890	4465
20	14	14544	15893

TABLE 3

S. pneumoniae - Putative coding regions of novel proteins not similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)
21	3	3359	2589
21	5	4802	4482
22	121	17099	17362
22	125	119467	19982
22	133	125540	23764
22	135	126388	126218
22	136	126382	23572
23	7	6655	6032
23	8	7132	6653
24	1	36	518
25	5	13009	2641
27	4	4819	4223
27	5	4789	4956
28	5	3017	1797
28	8	4272	3850
28	10	5028	4397
28	11	5746	5072
29	7	5596	4919
29	8	5039	5518
29	9	5595	8207
30	9	6511	6263
31	6	2664	2344
32	5	5203	5538
33	8	5327	4668
34	10	8024	7740
34	12	9360	8641
34	13	9667	9377

TABLE 3
S. pneumoniae - Putative coding regions of novel proteins not similar to known proteins

Contig	ORF	Start	Stop
ID	ID	(Int)	(Int)
34	18	13104	11902
35	11	9688	8588
35	12	11073	9670
36	2	334	1041
36	12	11120	10893
36	13	10993	11388
36	15	12172	14595
38	7	4269	4577
38	8	4480	5001
38	10	5517	5711
38	17	10732	11376
40	3	1728	3143
43	1	172	5
43	7	8884	8732
43	8	9568	9071
44	4	4831	6831
45	3	3204	3665
46	4	3875	3468
46	7	6074	7081
48	5	3196	3582
48	8	4579	4229
48	10	16342	15764
48	11	9323	8922
48	16	13042	12494
48	19	21979	21776
49	1	209	3

TABLE 3
S. pneumoniae - Putative coding regions of novel proteins not similar to known proteins

Contig	ORF	Start	Stop
ID	ID	(nt)	(nt)
50	4	3307	2672
51	5	3239	3598
52	111	12146	12883
54	7	5588	5187
54	8	6013	5459
54	9	6004	6210
54	16	17685	17506
55	9	110515	10123
55	12	11947	12141
56	3	935	1387
56	4	1496	1939
57	3	1624	2110
57	4	2100	2501
58	6	7541	7335
59	1	2	430
59	4	2416	2736
59	5	2734	3063
59	8	4743	5549
59	9	5459	5929
60	6	5741	6451
61	3	2395	1772
61	5	3316	3176
64	1	2722	2
66	2	1180	3147
66	8	9082	9495
67	3	1343	1182
69	2	1165	980

TABLE 3
S. pneumoniae - Putative coding regions of novel proteins not similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)
70	5	4059	3922
70	6	4215	4057
70	9	5268	5504
71	15	20351	121901
71	16	21859	122338
71	19	22204	127556
72	9	8458	8081
73	4	3815	4216
73	6	4214	4582
73	7	4369	4773
73	10	7183	6438
73	15	9452	9668
76	1	524	195
76	2	867	535
76	11	8602	9210
80	6	7924	8109
81	1	244	2
81	10	6631	8931
83	4	1872	1150
83	17	16810	16460
84	3	4464	2939
86	2	2147	1092
86	4	3606	2875
86	19	16767	17114
87	5	5326	5000
87	7	6459	6001
87	9	7224	7006

TABLE 3

S. pneumoniae - Putative coding regions of novel proteins not similar to known proteins

Contig	ORF	Start ID	Stop (nt)
ID	ID	(nt)	(nt)
87	18	17930	17670
87	19	18275	17928
88	2	1619	1840
88	4	2711	2878
88	9	6252	6016
89	3	2634	1621
89	9	7371	6868
90	2	899	2395
90	3	1143	952
91	3	2959	3141
91	4	3170	3691
91	6	4253	4573
93	1	391	2
93	6	2668	2379
93	8	4533	3712
96	1	3	182
96	2	904	632
96	3	1407	1147
96	4	1250	1420
97	9	7043	6753
99	15	18522	18592
99	17	19717	19341
100	2	4094	1980
103	1	48	239
103	6	4924	4373
104	5	6142	6735
105	7	6098	6517

TABLE 3

S. pneumoniae - Putative coding regions of novel proteins not similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)
106	1	1	363
106	10	9832	10212
108	1	2	268
111	3	3417	3788
111	4	3809	4606
115	10	10854	10438
116	3	2873	2121
118	2	2274	1357
122	4	2698	2333
122	10	5858	6139
122	12	6301	7416
124	2	346	690
128	4	2544	3368
129	1	689	102
129	2	1011	724
129	8	6454	6056
129	9	6540	6277
129	12	7809	7621
131	3	1433	756
131	10	5972	5673
134	11	11838	111209
135	2	625	1140
136	4	2913	3830
137	2	325	134
139	12	16027	14521
139	13	14840	14532
139	14	15363	14875

TABLE 3
S. pneumoniae - Putative coding regions of novel proteins not similar to known proteins

Contig	ORF ID	ORF ID (int)	Start (nt)	Stop (nt)
140	20	19822	20838	
142	1	1	1	285
146	3	760	479	
146	4	1149	778	
146	7	3604	2885	
146	13	8223	9401	
146	14	9399	10676	
146	15	110052	9750	
147	7	7488	7276	
147	9	8913	8647	
148	7	5298	4765	
149	1	2	1936	
149	3	2557	2880	
149	9	6258	6070	
150	2	1355	579	
150	3	2556	1909	
153	3	2061	2642	
154	3	1953	1741	
155	2	2181	1411	
156	8	4550	4311	
159	4	1384	1722	
157	1	37	294	
159	7	3271	4017	
159	12	631	780	
161	2	1332	1018	
165	3	5535	4945	
166	6	5406	4972	

TABLE 3
S. pneumoniae - Putative coding regions of novel proteins not similar to known proteins

Contig	ORF	Start	Stop
ID	ID	(nt)	(nt)
167	9	6075	6395
169	5	2928	3205
170	7	6485	6243
170	8	6964	6362
170	9	7303	6962
170	11	8790	7906
171	9	7150	7476
172	15	2298	1948
173	4	2913	2677
175	2	659	835
175	3	893	1789
176	2	1487	546
176	3	2200	1466
177	9	4686	4925
177	10	4923	5177
177	11	5111	5347
177	13	7396	8703
178	6	3452	3724
181	5	1853	2473
182	2	2112	1102
185	5	4683	4219
185	6	4846	4634
187	4	2940	3557
188	4	3686	4363
188	5	4183	4821

TABLE 3
S. pneumoniae - Putative coding regions of novel proteins not similar to known proteins

Contig ID	ORF ID	Start (nt)	Stop (nt)
188	6	5882	6493
189	5	3143	2844
189	9	5936	5364
191	1	618	4
191	11	10357	10001
192	3	2861	2268
192	4	3081	2878
192	7	6800	5331
193	3	997	839
194	4	2315	2127
195	5	6249	4543
195	6	6620	6231
196	2	1553	1849
197	1	1	861
198	9	6844	6644
200	5	5329	5769
200	6	5993	6595
204	5	3914	3276
205	2	447	1709
209	4	2038	2460
209	5	2458	2682
210	10	7370	8230
210	13	9029	10441
210	14	10439	10705
214	5	2581	2330
214	9	5065	5277
214	11	5996	5754

TABLE 3
S. pneumoniae - Putative coding regions of novel proteins not similar to known proteins

Contig	ORF	Start (nt)	Stop (nt)
ID	ID	(nt)	(nt)
217	2	541	194
218	2	914	1432
218	3	1430	1972
218	6	3639	3821
219	1	458	39
220	1	869	600
223	4	2617	1964
227	1	1	510
234	4	1539	1312
234	6	2116	1838
235	1	52	312
235	2	310	687
238	1	660	64
246	1	1	270
248	1	3	162
248	2	443	1222
254	3	2789	792
258	2	1179	1616
260	3	1770	2123
263	1	653	177
263	4	2244	1900
263	5	3569	2973
266	1	1	342
266	2	177	1022
270	2	1124	1681
272	1	857	186
275	2	1684	2295

TABLE 3 *S. pneumoniae* - Putative coding regions of novel proteins not similar to known proteins

Contig	ORF ID	Start (nt)	Stop (nt)
	ID	(nt)	(nt)
278	1	2	406
282	1	714	391
282	4	1463	1134
287	2	1119	826
288	1	540	4
289	1	684	4
291	5	1589	1858
293	2	2539	2925
294	1	21	608
296	2	494	700
296	3	670	843
302	1	261	530
309	3	559	350
310	2	249	1889
316	1	2087	1818
317	2	1048	584
318	2	313	777
319	3	477	133
327	2	912	607
331	1	1	549
333	1	2	535
333	2	465	82
333	3	127	342
341	1	1	705
345	2	895	701
346	1	750	199
349	1	1	198

TABLE 3
S. pneumoniae - Putative coding regions of novel proteins not similar to known proteins

Contig	ORF	Start	Stop
ID	ID	(nt)	(nt)
350	2	81	413
355	1	44	973
358	2	636	448
360	2	948	628
364	2	1639	1265
378	1	345	1004
379	2	683	510
381	1	109	693
385	1	150	4
385	2	269	30

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(D) SOFTWARE: ASCII Text

(vi) CURRENT APPLICATION DATA:

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(A) APPLICATION NUMBER:

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150

(2) INFORMATION FOR SEQ ID NO: 1:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 5625 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:

CCAAGCAAAA	CCAGCTACAG	CTAAAGGAAC	TTACGTAACA	AACTTGACTA	TCACAACTAC	60
TCAAGGTGTT	GGTATCAAAG	TTGACGTAAA	CTCACTTTAA	TCAGTAGTTA	AAGTAATGTA	120
AAAAAGTTGA	AGACGCTATG	TCTCAACTTT	TTTGATGTA	CGACGGGCAT	GTTGTATAGT	180
AGATGTGTAC	TATTCTAGTT	TCAATCTACT	ATAGTAGCTC	AGAAGTCGGT	ACTTAAACGT	240
GCTATATCAA	AACCAGTCCT	TGAAAAACGT	GGACTGGTTT	CGTGTGGGA	TTATTACCTT	300
GAACGACATG	CGTTAAAAGT	TAGTTGAACC	GCCGTATGCC	GAACGGACGT	ACGGTGGTGT	360
GAGAGGGGCT	AGAGATTATC	CCCTACTCGA	TTTCGAAATC	TAGTGGAAATG	AATCTGGAAT	420
AGTCCATCGA	GCTTTCTAAT	ACTCTTCGAA	AATCTCTCA	AACCACGTCA	ACGTCGCCTT	480
GCCGTGGTAA	TGGTTACTGA	CTTCGTCAGT	TCTATCCACA	ACCTCAAAAC	AGTGTGGTGA	540
GCTGACTACG	TCAGTTCAT	CTACAACCTC	AAAACAGTGT	TTTGAGCAAC	CTGCGGCTAG	600
TTTCCTAGTT	TGCTCTTGG	TTTCATTGA	GTATAACACA	TTGTTAGAAG	TTGGTTAAA	660
TTTCCTAATC	AGTTTGTCA	CATTTACCTT	CGATATATTA	TATCCCATAG	TTAAGGTTGG	720
TCATACAGAT	GATTATAGTC	ATGGAGCCGT	AAAACAGTGT	TTTCTTTAG	TTGACAAAGA	780
TGCCATGAAA	AAAATATTTG	TAACTGTAAT	AGGATATTT	GAAATAAATA	TAGATGAAAA	840
TATCACCGAT	ATTCTATACG	AAATGGTAC	TGCTATTCTT	TATCTTTATT	TACGTTCAAT	900
TGTTTCATAA	GTTCGGCAA	TTGATAGCAG	TGAAGCAATG	TTGCTACCTA	TCATTAATGT	960
TTTAGACTTA	CTAGATAAAT	CTCAACCTTT	TGAAGAAGAA	TAATTATTA	GCTCACTAJA	1020
TTGAGGGTAA	GGAAAAGTAA	AAGCAGTAAG	AAAATGTCT	TGCATTATAC	AGCAACCTTT	1080
TGGGAATGAG	TGGATGGATT	GAATAAAATT	TGATTAAGAG	TGGATGATTT	ATCTGTAGAT	1140
TATTATGGA	CAGTTAGTCT	TGAAGTAGTC	TAAGAATTAG	GTTATAATCA	GTAGAAGCCT	1200
TGCTAATAAT	GAGGAGGTTA	GTATTATGTAT	AGTAGACTGA	ATCTAAAATA	GTACGAAACA	1260
ATTGCTAAAA	CATTTATAGA	AAATTAATTT	ACTTTCCCAA	TCGATTGTTT	CTCATCTTAT	1320
TTCAATCCGC	TATATATTAT	GGTATCGAAT	CTTCATCAGA	ATGATAAAAT	TAATCAATTG	1380
ATATCTGATT	ACAAACAGAA	TATGAAAGCT	TTTTATATCA	CTATTGAAAA	ATTTATACGA	1440

151

GATGATGAAA	GCCTTAAGTG	TTATTTTATA	AAGGTTATTT	CAAGTCGTT	CAAGGTAACA	1500
AGTCTAGATC	AGATTGAAGC	TGATAAAAACG	ATACAAAGAA	AATATTCAAG	TGAGCTAAA	1560
AAATTTATTG	GATTTATAA	TGAGATTATT	TCTGAGGAAA	ATAGTTTCT	ACATGTACCA	1620
AAGAGGTGGT	CGAGTTGGTT	TAGGTTAGTCG	ATGCGTAGT	TGATAATTCT	CAGGGTATGG	1680
ACTTCTTTT	CATGAATGAG	GTAAAACAGC	AGGTATTGTT	TAGAGACAAT	CATTCTGAGC	1740
ATATTTCTG	GATAGAGGGA	GTATCCGATT	TTATGATCAA	AGTTAATACC	GCCCTCTGGT	1800
GAGAAGATGA	GTAGGTTGGT	AATTTAAACT	ATTAACAGA	ATTTTGATT	AAAAGTATTA	1860
TTTCATGAGA	GAAATCCTAA	TTTCACAATC	CATAGGCAA	CGCTTGCATT	TCGTTTTTA	1920
TTGGACTATA	ATAGGTTGGT	ATAAAGCCTT	CTGTAGTAAT	AAAATGTAGA	AGGTGTAGAA	1980
AGTAAGGATT	TAGAATATT	GTAGTTAAAA	ACACAATGTT	GCTATTCCCT	ACGATAGGGA	2040
GATAGATATG	GCAATGATAG	AACTGGAACA	TCTTCAGAAA	AATTTGTGA	AGACTGTTAA	2100
GGAAACCGGGC	TTGAAGGGGG	CTTTCGCTC	CTTTATTCTA	CCTGAAAAGC	AGACCTTTGA	2160
AGCGGTCAAG	GATTGACCT	TTGAGGTTCC	AAAAGGGCAG	ATTTTAGGAT	TTATCGGGC	2220
AAATGGTGT	GGGAAGTCGA	CAACCATTAA	AATGCTGACA	GGAATTGTTGA	AACCAACATC	2280
TGGTTTTGT	CGGATTAACG	GCAAGATTCC	CCAGGACAAT	CGGCAAGATT	ATGTCAAAGA	2340
TATTGGCGTA	GTCTTGGAC	AACGCACCCA	GCTATGGTGG	GATTGGCTC	TGCAAGAGAC	2400
CTACACTGTC	TTAAAAAGAGA	TTTATGATGT	GCCAGACTCG	CTCTTTCTATA	AGCCTATCCA	2460
CTTTTGAAT	GAAGTCTTGG	ATTGAAAGGA	CTTTATCAAG	GATCCCGTGC	GGACTCTTTC	2520
ACTGGACAA	CGGATGCGGG	CGGATATTGC	GGCCTCCTTG	CTCCACAATC	CCAAGGTTCT	2580
TTTTTTAGAT	GAGCCGACCA	TTGGTTGGA	CGTTTCGGTT	AAGGATAATA	TTCGTCGGGC	2640
AATTACTCG	ATCAATCAAG	AGGAAGAAC	TACCATTCCTT	TTGACCACTC	ACGATTTGAG	2700
TGATATTGAG	CAACTTGTG	ATCGGATTTT	CATGATTGAC	AAGGGGCAAG	AGATTTTGA	2760
TGGAACGGTG	AGCCAACCTCA	AGGAGACCTT	TGGTAAGATG	AAGACTCTCT	CTTTGAACT	2820
GCTACCAAGGT	CAAAGTCATC	TCGTCTCTCA	CTATGACGGT	CTGTCGATA	TGACCATGTA	2880
TAGACAAGGA	AACAGCCTCA	ACATTGAATT	TGATAGTTCT	CGCTACCAGT	CAGCTGACAT	2940
TATCAAGCAA	ACCCTGTCTG	ATTTGAAAT	CCGCGATTTG	AAGATGGTGG	ATACGGATAT	3000
TGAGGATATT	ATCCGTCGCT	TCTACCGAAA	GGAGCTCTAG	GATGATCAA	TTGTGGAGAC	3060
GTTATAAAC	CTTATCAAT	GCAGGGGTT	AGGAGTTGAT	TACTTACCGA	GTCAACTTTA	3120
TTCTCTATCG	GATTGGCGAT	CTCATGGGGG	CTTTTGTGGC	CTTTTATCTC	TGGAAGGCTG	3180

152	
TCTTTGATTC TTGCAAGAG TCTTTGATTC AGGGCTTCAG TATGGCGGAT ATCACCCCTCT	3240
ACATCATCAT GAGTTTGTG ACCAATCTTC TGACTAGATC CGATTCTGCC TTTATGATTG	3300
GGGAGGAGGT CAAGGATGGC TCCATTATCA TGCGTTGTT GCGACCGATG CATTTCGCG	3360
CCTCCTATCT TTTCACCGAG CTTGGTTCCA AGTGGTTGAT TTTTATCAGC GTTGGCCTTC	3420
CATTTTTAAG TGTCATTGTC TTGATGAAAAA TCATATCGGG TCAAGGTATT GTAGAGGTGC	3480
TAGGATTAAC TGTCATTTAT CTTTTTAGCT TAACGCTCGC CTATCTGATT AACTTTTCT	3540
TTAATATTTG CTTTGGATTT TCAGCCTTTCG TGTTTAAAAA TCTTTGGGGT TCCAACCTAC	3600
TTAAGACTTC CATACTGGCT TTTATGTCGG GGAGTTTGAT TCCCTTGGCA TTTTTTCAA	3660
AGGTTGTTTC AGATATTCTC TCCTTTTGC CTTTTTCATC CTTGATTTAT ACTCCAGTTA	3720
TGATCATTGT TGGAAAATAC GATGCCAGTC AGATTCTCA GGCACTCCTT TTGCAGTTCT	3780
TCTGGCTCTT AGTGTGGTG GGATTGTCTC AGTTAATTG GAAACGGGTC CAGTCCTTTA	3840
TCACCATTCA AGGAGGTTAG TATGAAAAAA TATCAACGAA TGCACTTGAT TTTTATCAGA	3900
CAATACATCA AACAAATCAT GGAATATAAG GTAGATTTG TGGTGGTGT CTTGGGAGTC	3960
TTTCTGACTC AAGGCTTGAA TCTCTGTTT CTCAATGTCA TCTTTCAACA TATTCCATTTC	4020
CTAGAAGGCT GGACCTTCA AGAGATAGCT TTCATTATG GATTTCCCTT GATTCCCAAG	4080
GGAAATGGACC ATCTCTTTTG TGACAATCTC TGGGCACTAG GGCAACGCCCT AGTCCGAAAA	4140
GGGGAGTTG ACAAGTATCT GACTCGTCCC ATCAATCTC TCTTTCACAT CCTAGTTGAA	4200
ACCTTTCAAGA TTGATGCCTT GGGTGAACCTC TTAGTCGGTG GTATTTTATT GGGAAACAACA	4260
GTGACCAGCA TTGTTGGAC TCTTCCAAAATC TTCTGCTTT TCCTAGTTG TATTCCCTTT	4320
GCGACCTTGA TTTATACTTC TCTTAAATCA GCAACAGCCA GTATCGCCTT TTGGACTAAG	4380
CAGTCAGGCG CCATGATTAA CATCTTCTAT ATGTTCAATG ACTTTGCTAA GTATCCGATT	4440
TCTATTACAA ATTCTCTTCTC TCGTTGGTG ATTAGCTTTA TCGTGCCTTT CGCCTTTACA	4500
GCCTACTATC CAGCTAGCTA TTTCTTACAG GAAAAGGATG TGTCTTTAA CGTAGGAGGT	4560
TTGATGTTGA TTCTCTGGT TTTCTTGTGTT ATTCCCTTA AACTTTGGGA TAAGGGCTTA	4620
GATTCCATCG AAAGTGGGGG TTGCTAAAG CTAAGAAG ACTAAATCA AGAAAGAAC	4680
TTATGATGTT TGTAATTGAA GAAGTCAAGG ATGAAAATCA AAAAAAGGCA GTTGTGCTG	4740
AGGTTTGAA GGATTGCCA GAATGGTTG GAATCCCAGA AAGCACACAA GCCTATATAG	4800
AAGGAACACAC GACACTGCAA GTTGGACCG CCTATCAGGA GAGTGTGTTG ACTAGATTTG	4860
TAAGCTTATC CTATTCGAGT GAAGATTGTG CAGAGATTGA TTGTCTCGGC GTAAAAAAGC	4920
TTATCAAGGT AGAAAATTC GGAGCCAATT GCTTGCTACT TTAGAGAGTG AAGCTCGTAA	4980

153

AAAAGTTGGT TATCTGCAGG TCAAAACAGT GGCAGAAGGT TCTAATAAAG ATTATGATCG	5040
AACAAATGAC TTTTATCGAG GTCTTGGCTT TAAAAAGTTA GAGATTTTC CTCAACTATG	5100
GAATCCGCAA AATCCTTGTC AGATTTGAT TAAAAAGCTT GAATAATATT ACTTGACATC	5160
TATTCTCAGA GTGCTATACT GAAAGTGTAA TCGCCGATTT AGCTTAGTTG GTAGAGCAAG	5220
GCACCTCGAA AGCCTAGTT ATAGGTAGAT AAACGACTGA GGATTTGAAA AAATAGATAG	5280
GTAGAAGATA ACCGTTAACG CTTACTCTTA GCGGTTATTT ATATTGTTTA ATAGCGCTAA	5340
TATTTTATCA ATTATGCCTG TTTTCGTGTT TCTGGTAGTT GTTCAAGTTT ATTGCTACTA	5400
TTTTGATGG TATGAATGTG CTTATAATGT ATCCCGTTA ACGAAAGTTT TGGACTTATA	5460
CTCTTCGAAA ATCTCTCAA ACCACGTCAA CGTCGCCTTG CCGTGCCTAT GGTTATGACT	5520
TCGTCAGTTC TATCCACAAAC CTCAAAACAG TGTGTTGAGT GACTACGTCA GTTCCATCTA	5580
CAACCTCAAAC ACACGTGTTT GCCCAATCTG CGGCTAGTTT CCTAG	5625

(2) INFORMATION FOR SEQ ID NO: 2:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 7571 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:

CTCTCCAGCT TTCCCTTGCAGA GTTGGCCATG TTGTGTCTTT AAGAAGTCTA AAAATATCTC	60
CAATAAAACG CATCGCTCTC TCCTATCTCG TTTCTCTGTG TGTAGTGTAC TTGCCACAAT	120
GCTTACAAAA TTTATTTACT TCTAGTCGTG TAGGCTTGAG GTTTCGCTG ATCTTGATTTG	180
AATAGTTTCT CGAACCAACAA ACCGCACAAAG CTAGGCCTTGC TTTTTTTAGT GCCATAACGC	240
CTCCATCTTA TCCATTATAA CAAGAAAGCT AGGCTTGAC AAGCATCTTA GCGAAATAGA	300
TTGACTATCG AATCCCATAT TGTTTGAGCC TTTTCCTTAA TCTTCGCATC TGAGATAGCC	360
CGGCTAGCCT CATCTACTAG ACTTTGCGCA CGCCCTCGAA TATCAGACAA ATTATCATCT	420
GTCTGGCTAT TATCATTGGT TTGTACTTGT CTTTTGTAT TGGCTGGTGC AATTCCATT	480
TGCTTATAAG CATTTCAAC CGTAAAGGTA CTTCCCTGGCG TATAAGGTAA AATGGTATTG	540
GCAATGTTTC TAAAGACATG AGCTGCACCG TTTGAAGTAG AGCCAGCTAG ATAGTGGTTT	600
TCATCAGTGG TCGGAAAGCC AAGCCAGTGG CTAATCACTA CATCCGGAGT ATAACCAATT	660
ACCCACTGGT CACTTGTGTA CTCCGGATTG AAAACTGCTT CAGTTGTTCC AGTTTCCCT	720

154

GCCATGACAT	AGTCTGCAGG	CGATGAACTA	ATACCGGTAC	CGTTGGTGA	AGTCCCCAAC	780
ATCATACTGG	TCATCTTGTC	AGCTACAGAC	TTATCAATCA	CCCGTTTTG	TGAATTTTA	840
TGACTCGAA	TAACCTGTCC	ACTAGCATT	TCAATTCTAC	TAATAAAATG	AGCTTCAGGC	900
ATTAAACCTT	CATTGCAAA	GGCGGCGTAT	GCTTGAGCCA	TTTGAAGAGG	GTTGGTTCA	960
ACACCGCTTC	CCAAGGCGAC	ACCAAGAAC	CGGTCGACCT	TTTCCATGTT	GAGTCCGAAT	1020
TTTTCGCCCTG	CCTCAAAAGC	CTTGTGAC	CCCAAATCAT	TAACAGTGGC	AACAGCAGGT	1080
AGATTAAGCG	ATTCTGCCAA	GGCTTGATAC	ATAGGAAC	CTCGACTCGT	TTTGATCCCT	1140
GCATAGTTAT	CAACCTTATA	GCTGTACATAC	TGCATGGTAT	GGTTATCCAA	CTGCTTATTC	1200
AAAGCCCAGC	TTGCTTCAAC	TGCTGGCGTA	TAACAACTA	AAGGCTTAAT	TGTAGAACCA	1260
GGACTACGCT	TTGATTGGGT	TGCATAGTG	AAATTCCGGA	ATCCAGTTTT	ATCATTGTCA	1320
GCAACTTGAC	CGACAAC	ACGAAC	CCCC	CCTGTTTCG	GTTCGAGGGC	1380
GATTGAGCAA	ACGTTCCATC	CTCTGCCCTC	GGAA	ATAGCG	ATGTTTTC	1440
TGCATATTG	CTTGGTAGTT	TTGGTCCAGC	TCTGTGAAA	TGCGGTAGCC	ATTATTGACA	1500
ATCTCTTCCT	CTGTTAGATT	ATAC	TTGGAA	ACAGCTCAT	TAACCACCGC	1560
GAGGGGTAAC	GGTAATCTGA	GATT	TTTCCT	TCATACTTAT	CGTCAATTG	1620
TCAACTTCAG	CAGCTTG	TTCTTGTTT	TTATCAATAT	ATCCTGCTGC	AACCATATT	1680
TGCAAGACAG	TATCGCGCCG	ATTAGTAGAA	TCTTCTACGG	AATTCAAGGG	ATTATACAGT	1740
TCCGGCCCT	TGACCATCCC	TGCGAGGTC	GCAGCTGAT	CCAGACTCAC	TTCTGATGCA	1800
GAAACTCCAA	AGTATTCTT	ACTCGCATCT	TCTACACCCC	ACACACCATT	TCCAAATAA	1860
GCCTGTTAA	GGTACATGGT	TAGAATTG	TCCCTACTAT	ATTTTTGCT	TAATTCTAAG	1920
GCAAGGAAA	ATTCTTCGC	TTTCTCTCA	ACAGTTGAT	CCTGCATAA	ATAGGC	1980
TTAGCCAGCT	GTGGGTAAT	GGTAGAGCCA	CCACCTGAAC	GTCCAGCAGT	GACAATAGCC	2040
AAGAAAAAAC	GGCCATAGTT	AATCCCGTCA	TTTTTATAGA	AAGAACGGTC	TTCTGTCGCA	2100
ATAACAGCAT	TCTGCAAGTT	TTTACTGATG	TCAGTCAGCT	CAACATAGGT	TCCCCTTTGA	2160
CCAGACAAGG	CACCAGCCTC	TTTTCTTCA	CGGTCAAAA	TAAGAGTCCG	AGTTTCAAG	2220
GCATTTGCA	AATCATTGAC	ATTGGTGCAC	TTGGCTACAG	CAAACAAATA	GATTCAACT	2280
ACCAAGCCTG	CACTCAAACC	TAGTATAAGG	ATAATCTTG	TTAGATGATA	ACGACGCCAG	2340
AATTTTCGAA	TCGGACCTAC	TTGGGCTAA	TTTTTCGAT	CACTACGAGA	GCGACGTAAG	2400
ATAGTGAAT	CAGAGTCCTC	TAGTTCACTT	TTTCTTTT	TAAAAAGAGA	AAGAAATTTC	2460
TCAAATAATT	TATCTAATT	CATGCGTTA	TTTATCATC	TTCATCATAG	GAAGACAAGA	2520

ATTTAGCTAT	TTCCTATCCA	AATAGGGCTT	TTTTGTTAC	AATATCTGTA	TGCAATTAC	2580
ATTTACATTA	CCCGCCTCTC	TACCTCAAAT	GACAGTAAAG	CAATTACTG	AGGAACAACT	2640
CCTCATCCCT	AGAAAATCC	GTCATTTTT	GAGAATCAAG	AAACATATTT	TGATAAAATCA	2700
AGAAGAAGTC	CACTGGAAAG	AAATCGTAA	TCCTGGAGAT	GTTCGCCAGT	TGACTTTG	2760
CGAGGAAGAT	TATCCCCAA	AGACGATCCC	TTGGGGCAAC	CCAGACTTAG	TGCAGGAAGT	2820
TTATCAAGAT	CAACACTG	TTATTGTAA	CAAACCAGAG	GGGATGAAAA	CGCATGGTAA	2880
TCAACCAAAC	GAAATTGCC	TTCTTAACCA	TGTCAGTACC	TATGTTGCC	AAACCTGCTA	2940
TGTCGTTCAT	CGTCTGGACA	TGAAACAG	TGGCTTAGTT	CTCTTGCCA	AAAATCCTT	3000
TATCCTGCC	ATTCTCAATC	GCTTATTGGA	GAAAAAAGAG	ATTTCTAGAG	AATATTGGC	3060
TCTAGTTGAT	GGAAATATCA	ACAGAAAAGA	ACTTGTTC	AGAGACAAAA	TTGGACGTGA	3120
TCGCCATGAT	CGTAGAAAAA	GAATAGTTGA	TGAAAAAAT	GGGCAATATG	CTGAAACGCA	3180
TGTAAGCAGA	TTAAAGCAAT	TCTAAACAA	GAATTCCTTG	GCTCATGCA	AGCTAAAGAC	3240
AGGGCGAAC	CATCAGATTC	GTGTGCACCT	TTCCGATCAT	AATCTTCTA	TCCTGGGAGA	3300
CCCTCTCTAT	AATAGTAAAT	CAAAGACAAG	CCGGCTTATG	CTTCATGCC	TCCGACTTT	3360
CTTTACCCAC	CCACTTACTT	TAGAGAAGCT	AACTTTCACT	ACCCCTTCAA	ATACATTG	3420
AAAAGAATTA	AAAAGAATG	GATGATCGTG	TCATCCATT	TTCCATATAA	AAAAGCAAGA	3480
CCACAAAGCC	TTGCTTTCTA	TCAACTCAAG	AATTATTTAG	CAATTTTTC	GAAGTATTCA	3540
AGAGTACGAA	CAAGTTGTGC	AGTGTATGAC	ATTCGTTGT	CGTACCATGA	TACAACCTTA	3600
ACCAATTGTT	TACCGTCAAC	GTCAAGAACT	TTAGTTGAG	TTGCGTCAA	CAATGAACCG	3660
TAAGACATAC	CTACGATATC	TGAAGATACG	ATTGGATCTT	CTGTGTAACC	GTATGATTG	3720
TTTGAAGCTG	CTTTCATAGC	TGGCTTCACT	TCATCAACAG	TAACGTTCTT	TTCAAGAACT	3780
GCTACCAATT	CAGTAACCTGA	TCCAGTTGGA	GTTGGAACGC	GTGTGTCAGA	TCCGTCAAGT	3840
TTACCATTCA	ATTCTGGGAT	TACAAGACCG	ATAGCTTTG	CAGCACCACT	TGAGTTAGGA	3900
ACGATGTTG	CAGCACCACT	GCGAGCACGG	CGAAGGTCAC	CACCAACGGT	TGGTCCGTCA	3960
AGGATCATTT	GGTCACCACT	GTAAGCGTGG	ATAGTAGTCA	TCAATCCTTC	AAACACACCA	4020
AAGTTGTCTT	GAAGAGCTTT	AGCCATTGGA	GCCAAAGCAGT	TTGTAGTACA	TGAAGCACCT	4080
GAGATAACTG	TTTCAGTACC	GTCAAGAACG	TCGTGGTTAG	TGTTGAATAC	AACTGTTTA	4140
ACGTCGTTTC	CACCAAGGAGC	AGTGATAACA	ACTTTTTAG	CTCCACCTT	AAGGTGTTT	4200
TCAGCTGCTT	CTTTCTTAGC	AAAGAAACCA	GTAGCTTCAA	GAACGATTT	TACACCGTCA	4260

156

GTAGCCCCAGT CGATTGTTC TGGATCACGT TCAGCAGAAA CTTTGATGAA TTTACCGTTA	4320
ACTTCAAAATC CACCTCTTT AACTCAACA GTACCGTCGA AACGACCTTG AGTTGTGTCCG	4380
TATTTCAACA AGTGTGCAAG CATAACTGGA TCTGTAAAGGT CGTTGATGCG TGTAACCTCA	4440
ACACCTCTA CGTTTGGAT ACGACGGAA GCAAGACGAC CGATACGTCC GAAACCGTTA	4500
ATACCAACTT TAACTACCAT TAGTGATTT TCCTCTATGA AAATCATGAA ATTTTTATTG	4560
TGAAAAGAGT AACTTGAATC ACTACAAATC ACCTTCAAC AACCTATTAA TACAACATT	4620
TGAGTTGAAT TGCAAGTATG GCCATTGTTT TTCTATGTTA GTTTCTTTT AAGACTGTAA	4680
ACCAAGGAAT CCCTTACTAT TCATAGCATA ACGATTCTAT AGGATCCATT TTACTAATCT	4740
TACCGGCCGG GAAGTAGGCT GAGACATAAC CAAGTAATAG AGCGAAAAGT AGAGTTCCCTA	4800
AAACAGATAA AAGATTTAAT TTAAAAACCT TAGTGATGGA TGGGTAAG TGACTTACAA	4860
TCGCATTCGC CAAACCTCCC ACCCCTTGTG CAACCAAAA TGCCAGCAGC AAGGGCATGC	4920
CTACAATCCA GATAGCCTCG TAAATAAAAA TTCCCTTGAC ATCACGATTC TGATAACCAA	4980
CTGCTTTCAT GACACCTATT TCCTTGGAAC GTTGATGAT ATTGATGTAATAATGATAC	5040
CAATCATAAC CGCTGCTACC ACAATAGCTT GTGATGAAAG CACAATCAAT AATCCCTGAA	5100
TAACACGAAT AAAGGTAATC ACAATATCAA GAACTCTCTG TTGAGAAAGC ACAGTATACT	5160
TCTTATTTT CTGTAATTCT TCTGTTACTA CTTTTGTCTG TGATGGATCT TTGAGTTCCA	5220
AGATAAAAATA AGATACAGCT TTGTAATC CAGCCTCTTT CAAAATCGTT TCCATTTGAT	5280
GAGACAGCAT GAAACTGTTG CTGCTCTCCA TGTCACTTC ATCACGATTC ACACGTACAA	5340
TCTTCCTTG AAATTGAGCA ATCTTAATAG TTTGGCAGC ACTTTCTACA ATGCTGGCTG	5400
AGACTGATTT GCCAATAAGA TCATTAAGCTG TCAAATTTT TCCTGTCTGT TCATTCCAAT	5460
TTTTTAGTAA ACTGCTTGGA ATCGTTAATC CCTGTTCAATT TGTATCAGTA TAGAGGGATC	5520
CAGCCAACAC TTTGTCGTC TCATTATTAC TAACAGAGAT ACTTGTATCA TCATAAAGAC	5580
TCACTACTTG AGCATAAGAA GGCATCGTTT GACTCAGATC CATTCTTGC CCATCTATAG	5640
TAATATTGTA CATGTTCATC CAAAAGGAC TCTCCAAATA TTTAATAGCT TCTTCCCAA	5700
CTGTATCCGT GATATATAGT CAATTGAAAC AAGAGCAGGA TAAAAAAGCC TCGTAAAGG	5760
TATTGCAACT TGTTAATACC TTTTGAGGT GCTTTTGAT ATGAGCCAT GTTTCTCAA	5820
TAGGATTTGTA CTCAGGGAG TAGGGAGGAA GAGGTAAAG TTTATGCCA AACTCTTCG	5880
ATAAAAGTTC TAGCTTCCCC ATTCTATGGA ATCTTACATT ATCCATAATA ATAACCGATG	5940
GTGTGTTAA TGTTGGTAAG AGAAAATTCT GAAACCAAGC TTCAAAAAG TCGCTCGTCA	6000
TCGTCTCTTC GTAAGTCATT GGAGCGATTA ATTCAACATT TGTAGACCT GCAACCAAAG	6060

157

AAATCCTCTG ATATCTTCTT CCAGATACTT TGCCTCTTAT TAATTGACCT TTTAATGAGC	6120
GACCATATTC TCGATAAAAA TAAGTATCGA ATCCTGTTTC GTCAATCTAA ACAGGTGCTA	6180
GGTGCCTTAA ACTATTTAAA TTCTTAAGAA ATAAGGTCAC TTTTCTGGG TCTTGTTCAT	6240
AGTAGGTGTG GTTCTTTTT CGAGTGTAGC CCATAGCTT GAGCGTATAG TGGATGGTAG	6300
TTGGATGACA GCCAAATTCA GAAGCTATTT CAGTCAAATA AGCGTCTGGA TTGTCAGTAA	6360
GATAGTTTT AAGTCTATCT CTATCAACCT TTCTTGGTT TATTCCCTTT ACTTGGTGGT	6420
TTAGCTCTCC TGTCTCTCT TTTAGCTTTA ACCAGCCATA AATGGTATTA CGTGAGATT	6480
GGAAAACGTG TGATGCTCT GTTATACTAC CTGTTCGCTC ACAATAAGAG AGAACCTTTT	6540
TACGAAAATC TATTGAATAT GCCATAAAAA GATTATACCA CATTGTGTAC TATTTTTGGT	6600
TCATTTTACT ATATTGAAAG AGGCCTTAA ACTATCTGAC ATAAAACCTCG TTCTAGAGGA	6660
AAGACATCCT TTAAAAAGTT AGTTTATTTT ACAACTAGA CATCAAGGTA GGTTAACCCC	6720
TTCATGGAAA AATCAAGACT CTTAGCACTA TGGGTTAAC TACCACTGGA GACGTAATCA	6780
ATCGCTAAAC CACGAAAACG GCTAATAGTG GTCATATCAA TATTTCCAGA ACATTCAATC	6840
CGAGAACGTC CTGCAATTAG GGTAAATGCC TGTTCAATCT GTTCCAATGA CATATTATCC	6900
AACATGATAA TATCAGCACC CGCCGCCGCA GCTTCTCGG CAGCAGCAAG GCTTCCACT	6960
TCCACCTCGA CCATTTTAC AAAAGGGCA TAGGCACCGG CTTGAGCAAT TGCCCTTTGA	7020
ACACTACCTA CTGCCGCAAT GTGATTGTCT TTTAGCAGGA TAGCATCTGA TAAATTAAAG	7080
CGATGATTAT AGCCACCGCC AACTCTCAGC GCATATTCT CAAAAGACG TAAATTAGGA	7140
GTAGTTTTTC GAGTATCAA TACCTTAATG CAATCATCGC CTAAGGCTTC TACATAAGCA	7200
GCTGTCATCG AAGCAATCCC TGATAATGT TGAAAAAT TCAAGGCAAC GCGTTCACAT	7260
GTAAAGAGAC TTCTCACCGA GCCTATGATT TCTAAAACCA AATGCCACT AGTCAAACGA	7320
TCCCCATCCT TAAATTGATG AGGATTCTGG AAGGTCACCT CGGCATCAA TAGGGTAAA	7380
ACCCTTGAA AAACGGTTAG CCCCGCTAAA ACACCAGCTT CCTGGCAA AAGCGACACC	7440
TTGGCTTGGC CATGATGATC AAAAATGGCA TTGGTACTGT AATCTTCGGA ATGAACATCT	7500
TCTCGCAAGG CTGCTTCAA TGTATCATCT ATTTGAAAAG GGGTTAAATC AGTTGAAATG	7560
ATTGACATCA C	7571

(2) INFORMATION FOR SEQ ID NO: 3:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 26385 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double

158

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:

TTTGCTAGTG	GCTTAAATTC	TTCAGGAAAA	TCAGCGTAT	CTAAAAGTCG	TGTCGTTTT	60
GTTTCATCTA	TATAAAGACT	TCCTGCTCCC	CCTACAACTA	GAAAACGTGT	CTGTGTTCCA	120
GCAAGAAGCT	GATTAATAG	TTCGATTGAT	TTGCTGTGGA	GCGGTAGCGT	ATCTGGTGT	180
TAAGCACCAA	ACGCTGAAAT	AACAGCATCA	AATCCAGTAA	GATCATCTTT	TGTCAACTCA	240
AATAAAATCTT	TTTTAATAAT	AGACTCAGCT	TGACTTTTGT	TTTCAGAACG	AACAATAGCC	300
GTTACTTCAT	GTCCTCGTTT	GACTGCTCT	TCAACAATTG	CTTTCCCCGC	TTGTCCATT	360
GCTGCAATAA	CTGCTAGTTT	CATTTTTAT	ACCTCTCTG	TTGTAATTAT	TTTAGTTACA	420
GAAATTGTGA	CACTCTTAAT	AATCAATGTC	AATAGTCTTG	CTTAATTATT	ATCAAATAT	480
TTCTACCAAG	AAAACTAACC	ATGATTCTAG	TGAAAAAAA	TCTTCTTTGT	CAACAAATT	540
ACTTTCTTGT	TTTAAACATG	CTATAATAAT	CATAGCAAGA	GATCTAAGTT	GTCTGTTT	600
TTAAAACGAG	GTGATTATCA	TGCGTAGATT	CTATTCCCAT	CTCCCCTACT	ATCTGGTCAT	660
ATTATTCTTT	TATTGCCAC	TTTATGAGTT	GTTCTTACTA	GTTGTTCTG	ACCCCTTAC	720
ACTCAAGGGA	CTCTATATAA	ACAATCTCT	CTTCTTTACA	CCTCTGGTAA	TCTTGATTGT	780
ATCGTTACTC	TATAGCTACC	GTTTCCGTTT	CTCACTTTGA	TGGTTAGTTG	GTAACGGACT	840
GCTCTTTAC	TTTACTATCA	TAACCTTTGG	TGAGTTATA	CTAATTACT	TGCTAATCTA	900
TGAAACAGTT	GCTCTGGTCG	GCATGGATTC	TGGTATTAGC	ATCAAGCATA	TTCTACAAA	960
AATGAAAAAC	AAAAACACTT	CACAAATCC	TTGAAAATC	TCACAATCAT	GCTATAATAA	1020
TCCATAGAGA	CAAGTCACCT	AGTCCCTTTC	TACTAGAGAG	TGCGTGGTTG	CTGGAAACCG	1080
ATAGGAAGTC	TAAACTGATA	CTACTCTTGA	GTTTTTTATG	AAAACATAAA	ACGGTGGCCA	1140
CGTTAGAGCC	GATCAGAGGT	GTCCCTCTCT	TTTGAGGTAC	ATAAAATGAAG	GTGGAACCAC	1200
GTTGCGACGT	CCTTCGAGG	ATGTCGCATT	TTTTTATTAG	GATACTAATT	ATGGAGTTGC	1260
AAGAATTAGT	GGAGCGCAGT	TGGCAATCC	GACAAGCTTA	TCACGAACTG	GAAGTTAAC	1320
ATCATGATTC	CAAGTGGACG	GTAGAAGAAG	ACCTCTTGGC	TTTATCTAAT	GATATTGGAA	1380
ATTTCCAACG	ACTGGTGATG	ACAAAGCAAG	GACGCTACTA	TGATGAAACA	CCCTACACAC	1440
TGGAACAAA	ACTTCAGAA	AATATCTGGT	GGCTATTAGA	ACTTTCTCAA	CGTTTGGATA	1500
TAGACATTCT	GACGGAAATG	GAAAACCTCC	TCTCTGATAA	AGAAAAGCAA	TTGAACGTTA	1560
GGACTTGGAA	GTAGTCTGCT	GATAAAAAT	CAATGCTTAG	AAACTATGAA	ATAATAAAA	1620

159

AGGAGAACAT CATGATTAAC ATTACTTTCC CAGATGGCGC TGTTCGTGAA TTGAAATCTG	1680
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CTGGTAAATT CAACGGCAA CTCATCGACA CTACTCGCGC TATCACTGAA GATGGAAGCA	1800
TCGAAATTGT GACACCTGAT CACGAAGATG CCCTTCCAAT CTTGCGTCAC TCAGCAGCTC	1860
ACTTGTTCGC CCAAGCAGCT CGTCTGTCTTT TCCCAGACAT TCACTTGGGA GTTGGTCCAG	1920
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ACCTTCCTCG TATCGAAGAA GAAATGCAA AAATCGTCAA AGAAAATTC CCATCTATTC	2040
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AATTGATTGA AGAACACTCA GAAGACGAAG GCGGTTTGAC TATCTATCGT CAGGGTGAAT	2160
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TCTACGGTAC AGCTTGGTTT GACAAGAAAAG ACTTGAAAAAA CTACCTTCAA ATGCGTGAAG	2340
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GGGCCCTTCCC AACATGGCTG GCACCACACC AAGTAACCCCT CATCCCAGTA TCTAACGAAAG	3300
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160

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TTCCCTTACCA ATTAATTGTT GGAGACAAAG AAATGGAAGA CGAAACAGTC AACGTTCGTC	3480
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162

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164

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165

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166

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ATCATCCAAA	ATAAGATTGC	AAGTGGATTA	TCAGAAGGAT	CTGTGAAGT	GACGAAAGAA	15180
GGAAAAAAAT	GAATAAAAGA	GGTCTTTATT	CAAAACTAGG	AATTTCCGTT	GTAGGCATTA	15240
GTCTTTAAT	GGGAGTCCCC	ACTTTGATTC	ATGCGAATGA	ATTAACATAT	GGTCAACTGT	15300
CCATATCTCC	TATTTTCAA	GGAGGTTCAT	ATCAACTGAA	CAATAAGAGT	ATAGATATCA	15360
GCTCTTGTGTT	ATTAGATAAA	TTGTCTGGAG	AGAGTCAGAC	AGTAGTAATG	AAATTAAAG	15420
CAGATAAAC	AAACTCTCTT	CAAGCTTGT	TTGGCCTATC	TAATAGTAAA	GCAGGCTTTA	15480
AAAATAATT	CTTTCAATT	TTCATGAGAG	ATTCTGGTGA	GATAGGTGTA	GAAATAAGAG	15540
ACGCCAAAA	GGGAATAAT	TATTTATTT	CCAGACCAGC	TTCATATTGG	GGAAAACATA	15600
AAGGACAGGC	AGTTGAAAAT	ACACTAGTAT	TTGTATCTGA	TTCTAAAGAT	AAAACATACA	15660
CAATGTATGT	TAATGGAATA	GAAGTGGTCT	CTGAAACAGT	TGATACATTT	TTGCCAATT	15720
CAAATATAAA	TGGTATAGAT	AAGGCAACAC	TAGGAGCTGT	TAATCGTGAA	GGTAAGGAAC	15780

167

ATTACCTCGC AAAAGGAAGT ATPGATGAAA TCAGTCTATT TAACAAAGCA ATTAGTGATC	15840
AGGAAGTTTC AACTATTCCC TTGTCAAATC CATTTCAGTT AATTTTCCAA TCAGGAGATT	15900
CTACTCAAGC TAACTATTTT AGAATACCGA CACTATATAC ATTAAGTAGT GGAAGAGTTC	15960
TATCAAGTAT TGATGCACGT TATGGTGGGA CTCATGATTC TAAAAGTAAG ATTAATATTG	16020
CCACTTCTTA TAGTGATGAT AATGGGAAAA CGTGGAGTGA GCCAATTGTT GCTATGAAGT	16080
TTAATGACTA TGAGGAGCAG TTAGTTACT GGCCCACGAGA TAATAAATTA AAGAATAGTC	16140
AAATTAGTGG AAGTGCTTC TTCATAGATT CATCCATTGT TGAAGATAAA AAATCTGGGA	16200
AAACGATATT ACTAGCTGAT GTTATGCCTG CGGGTATTGG AAATAATAAT GCAAATAAAG	16260
CCGACTCAGG TTTTAAAGAA ATAAATGGTC ATTATTATTT AAAACTAAAG AAGAATGGAG	16320
ATAACGATTT CCGTTATACA GTTAGAGAAA ATGGTGTCTG TTATAATGAA ACAACTAATA	16380
AACCTACAAA TTATACTATA AATGATAAGT ATGAAGTTTT GGAGGGAGGA AAGTCTTTAA	16440
CAGTCGAACA ATATTGGTT GATTTGATA GTGGCTCTT AAGAGAAAGG CATAATGGAA	16500
AACAGGTTCC TATGAATGTT TTCTACAAAG ATTGTTATT TAAAGTGAAT CCTACTAATT	16560
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GCCGCAAGGG AGGCCAATTA GTTGTGGTT TAGTCATAA AGAAGATGAT AGTATTGATT	17100
GGAAATACCA CTATGATATT GATTTGCCCT CGTATGGTTA TGCCTATTCT GCGATTACAG	17160
AATTGCCAAA TCATCACATA GGTGTACTGT TTGAAAATA TGATTCGTGG TCGAGAAATG	17220
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168

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AACTGAAATC TATTATACTA CAAACTATTG AAAGCGTTT AATTTTAAGG TATAATAATC	18540
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CCTAACATCAGC TTTTGATGGC TGATACTAGT ATCTTCGAAG AAGGGCTAGC AGCTGTAGAA	19020
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GACGGTCCAG ATTTGAATT GATTAAGAAA CCTCTGTGATG CTGGTGTAGA TGTCATTGCA	19140
GAAGGAAAAA TTCATACACC AGAACAAAGCC AAACAAATCC TTGAATATGG AGTGCAGGC	19200
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TCTGAATCCG CAGATTCAAC TTCAAATACT GCAGATTATT	20580
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170

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 GTCTTACAA AATCTCTGAT TAACACAGTT ATTTGGTTA TTGGATCTGT ACCAGTTGTT 22740
 GTTCTATTCT CACTCTTGT AGCATCTCAG ACCTATCATC AAAATGTCAT TGCCAGATCC 22800
 TTCTACCGTT TCGTCTCTT CCTTCCTGTT GTAACGGGTA GTGTTGCCGT GACAGTTGTT 22860

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TTTTGGAAGA	TTAAATGGCC	AAGCCTCTT	CCAACAACTC	TTTATATTGC	AATCATCACA	23160
ACAATTAACT	CATTCAGTG	TTTCGCCCTTG	ATTCAGCTTT	TGACATCTGG	TGGTCCAAAC	23220
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CAATTAAAG	TACTTGGAAA	CGACGTAGAA	TACTAAAGAA	AGGAGACAGC	TATGCAATCT	23400
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ACTGTGCTGT	TCATCTTCC	ATTCTACTGG	ATTTTGACAG	GGGCATTCAA	ATCACAACCT	23520
GATACAATTG	TTATTCCCTCC	TCAGTGGTTC	CCTAAATGC	CAACCATGGA	AAACTTCCAA	23580
CAACTCATGG	TGCAGAACCC	TGCCCCGCAA	TGGATGTGGA	ACTCACTATT	TATCTCATTG	23640
GTAACCATGT	TCTTAGTTG	TGCAACCTCA	TCTCTAGCAG	GTTATGTATT	GGCTAAAAAA	23700
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TATCAGAAAC	GAAGGAAAGA	GTATGATTT	TGACGATTG	AAAAACATCA	CCTTTTACAA	24540
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172

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TGTCCCTCAAT CAAGCTGAAA ATGATCAATT TGAGTATCAT AAGAACTATG CAGATTTGCA	24720
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AGCATTGAC GAAGCGAGTG ACATGGCTT TGTTCATTGT CATGAACACT ACCCACTCTT	24840
GTTGGGTTAT CACAATTGG CGATTTCTT CCCAGGTGAG CCACATCAGC CAAATGGTTA	24900
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CCTCATGTTA GCCATTCTTG TCCTCATTGT GATGGTTCTT TATCTGTCCG CCTTCAGTCT	26220
ACTCTTAGGT GGCTCAGTCT TCCTACTTTT TGGGTTGGA CTATTGGTCT TTATCCAGAC	26280
TGGATTGATG GAGAAAATT TCGAAAATA CCAATAGGAG CTTTTATTCT GAAACTACTT	26340
TCAAAGGCTC CAAACGCTAT TCTATAAGCG AGAAACTAAA ATCGG	26385

(2) INFORMATION FOR SEQ ID NO: 4:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 2716 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:

CCTGCCCGCA TTGCCCTAGG CATTAAAGTAA ACATATAAAA GCATGTGAGA GACTGTTGGA	60
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AAAGAAAAATG ATATAATAGT AGTTATGGAG AAAAGAAAT TACGCATCAA TATGTTGAGT	180
TCAAGTGAGA AAGTAGCAGG ACAGGGAGTT TCAGGTGCTT ACCGTGAATT AGTCGTCTT	240
CTTCACCGTG CTGCCAAGGA CCAATTGATT GTTACAGAAA ATCTTCCAAT CGAGGCAGAT	300
GTGACTCACT TTCATACGAT TGATTTCCC TATTATTTAT CAACCTTCCA AAAGAAACGC	360
TCAGGGAGAA AGATTGGCTA TGTGCATTT TCAGGCAGCTA CACTTGAGGG AAGTTTGAAA	420
ATTCCATTTT TCTTAAAGGG AATTGTGAAA CGCTATGTAT TTTCTTTTA CAACCGGATG	480
GAGCACTTGG TTGTGGTCAA TCCTATGTTT ATTGAGGATT TGGTAGCAGC TGGTATTCCA	540
CGTAAAAAAG TGACCTATAT TCCTAACTTT GTCAACAAGG AAAATGGCA TCCTCTACCA	600
CAAGAAAGAGG TAGTCAGACT GCGCACAGAT CTTGGTCTTA GTGACAATCA GTTTATCGTA	660
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ATGTTGCGTG ATTTAGATCT CTATAAGGTG ATTTTGGAGG GAAATTATCG GCGACAGCG	1020
GGTAGAGAAG AGATGAAAGA GGCTATTTG GAATATCAAG CAAATCCTGC TGTCTAAAAA	1080
GATCTCAAAG AAAAGGCTAA GAATATTTCC AGAGAGTATT CTGAAGAGCA TCTGTTACAA	1140
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174	
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CTAGAGATTCA AAGACTCAGT CATTTTACA GGGATGATTG CTCCTAGTGA GACGGCTCTT	2040
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TTATGGATTA TCGCGA	2716

(2) INFORMATION FOR SEQ ID NO: 5:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 13926 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:

CTTTGGTTTT GCCTTATTCA AGACATGAGG CCCATCAGGA ATGATCTGAA ACTGCGAATC	60
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TGTTAACAGT CTATGGAGAG CTTTCATAGA ACTAAGATTG	GGTTTATCTT TGCTGCCACA	120
AATTAGTAAG GTTGGATAAG GGTAAAGTTCC TGCTATATCC	GTAAATCAA GTGTCTTCAA	180
CTCCTCAGAA ACTCCGACCA TAAGAGTCTT GTCTGCTCCC	TGTTTTCAA ATACTCTTT	240
GGGAAGTAGT TTAAAAATCA GCAATTGAAG ATAAAATAGG	ATATTCCCTG CTAATTTAAG	300
CGGGCATCCT GACAGAATCA AAGCTCGAAG ATTTGGTAA	TCGTAACTGG AAAGTTCTAG	360
TGTCAGGGCA GCACCTAAGG ACAATCCAAT CAAAACAAAA	GGTTCTGTCT CTTGAGCTAG	420
GTGCTGATAA ACTCGCTCTT TAGCTTGTG ATAGTTACTA	ACTCCAGAAG GAAATAACTC	480
GATAGCCTCA GAAGGATAAT CTGTCAGTAG ATTCCGAACT	TCTTCCAAG ACTCTGCTGA	540
CTGCCCTAAC CCATGCAAAA ATATTAATTT CATCTAGTTC	TCCTCAAGGC TTAATTCTATA	600
CAAGCCTCTC ACTGCATTAC AGCCGTAAT AGCTTCTGCT	TGGGTTAAAT CTGCCAAGGT	660
CAAGACTTTC TCTTCTACCT GTCCCTGTTT TAGCAAATGC	TGACGGTAA TTCCTGGCAA	720
GATTCCAAGT CGGATAGGCG GTGTGTAGAG TTTTCAGCG	ATTTTCAGAA CCAAATTCC	780
TATAGAGGTT TCAAGCAGTT CTCCCTGACTT ATTGTGGTAA	ATCTTCTCTT GTTCTCCTAG	840
GCTCAAATGC GGTCGGTGAG TGGTTTAAA GTAGGTAAAG	GATTGATTCA AAGCAGCTTC	900
CTGAAGACAG ACTTGGGCCT GACAAAAGCT TGTACTGAGA	GGGGTTAATA CTTGACGATT	960
GACTTCTATC TCTCCAGATT TGCTAAGGCT GATTGCAAG	CGGTAATCTC GATTAGCTTC	1020
ACAATCCTGA CACTCTTCCT CAATCTTGTG TCCCAGTCT	TCTGCATCAA AACGAAAGC	1080
AAAATAACGA CTAGCTTTTC TCAGCCTTTC CAGATGTTGT	TCTTCAAACA TCAGTTGTT	1140
TTGGCTGATT TTTCCAGTTG TAATTAATTG GAAGCGAGCT	TGTTTACGAT AGAGAACTGC	1200
TGCCTTTGA TGAACCTCTC GGTATTTCAGA TTCCCATGTG	CTATCCCAAG TAATCCCTCC	1260
GCCAACCTCCA TAAATGGCTT GACCTTTGTG AAGTTGAATG	GTACGAATGG CCACATTAAA	1320
AATCCGTCGT CCATTGGAA GCAAGAGACC AATCGTTCCA	CAGTAGACTC CACGCCGTTG	1380
AGGCTCCAAG TCCTTGATAA TCTCCATTGT CGCAATTTC	GGTGCACCCG TTATGGAACC	1440
ACAAGGAAAG AGTGAGCGGA AGATTTCAAC AAGGCCACA	TCCCTCTCGCA ACTGACTCTT	1500
GATGGTCGAA GTCATCTGCC AAACAGTTGA ATACTGCTCT	ACCTGACACA GACGCTCCAC	1560
GTGCTCGCTC CCAACTTCAG AAATACGGTT CATATCATG	CGCAAGAGGT CCACAATCAT	1620
CATATTTCA GAGGGATTTT TGGGATCCTG TTCCAACCAA	CTGGCCTGTT CAAGATCTTC	1680
TTGGTCAGTT ACCCCACGCT GAGTCGTCCC CTTCATTGGT	CGTGTGTCAC ACTCGCGATC	1740
ATTTTGCTCA AAAAAGAGCT CTGGGCTCAT GGAAATCACT	GTCATCTCGT CATGTTCCAC	1800

176	
ATAGGCATTG TAGCCGCCT CCTGCTCTAC CACCATACGA TTGTAGATGG CAAAAGGATT	1860
GGCATTAACTTTGCTTAA GTTGGACGGT GTAGTTGACC TGATAGGTAT CTCCCTGCCG	1920
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CCAATTGAG GGCAAATCAA TATCCTCATA AGTCAGAGGA ATAGGGGAAG TTTCTACGAT	2040
ATCATGAACA GTAAAGTAAA GCAGGTACTC TCCCAGTAGG GGATCCTTGT GAACTGCTAA	2100
TTTTCTCTCA AAACGAGGTG CAGCCTCGTA GCTGACATAC CCCACCACAT AATAACCTTG	2160
CTCTTGGTAG CTTTCCACTT GTGCCAGCAA ATCTGCCACT TCTTCTACAT TTCTCGTTT	2220
CAACTCTTTA ATAGGCTGGG TAAAGGTATA TCTCTCCCCC AAAGTCTAA AATCAATCAC	2280
TGTTTTCTA TGCAACCTT AAGTATAGCA TAAAATAAGA AAACCCCTCAT CCGCAAAGCA	2340
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TAAATCATCG TATGATGAAA CGGAAGAACC ATTTACTTCG AATGTTGTTA ATCCTTTCGT	2580
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GAACCATGCA CTATTTAAAT CTTTATTGTTG TTGAGAAATC ACAGATTCA CCTCAATTTC	3120
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CATACCCCTGA GCATATGCTT CAGCATCAGT ACCTTCACGG TGTCAGAAC CACCTAAGTA	3360
AATCCGACGG TCGTTGACGT GTGTTGTTTC ATGTGTGTAAC TGAAATAC CGTATTCAACC	3420
AACCAATTCT AAATGAACAT ATTTTACATC AGTTCTAATA TCATCAGAGT TAGGATATAT	3480
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GGCAATTGCA	TTATAATTG	AAATTAAATA	AAGATGTGCT	TTATCAATAT	TCAGTAGTGG	3840
GAGTATAGTA	TTTCTAAGGT	GACTTCGTTT	TAAATTATCG	AATGCACGAT	GTTTAGAATT	3900
TTTAATTCT	TCGACCTCAG	AAGCGCGTTC	TGCGATGTAG	ACATGGCTT	CTGTAGCATC	3960
AATAAACCAA	TCGTTCATAT	TGTCTATATT	TGTGAACAAT	TGTCTATTAT	AATTAAAAA	4020
TGCATCTAAA	TTACCTGATT	TAGTATATT	AGCCAATACT	TGACCGAATG	CGTCGAATGT	4080
ACGTGAACCT	TTAATGTTGT	TCTCTTACA	ACCGATTCA	ATTAATCTGT	CTAATACGCT	4140
AACTTTTCA	CCATAGAAAT	CTGGTTGAA	TAGCATTAAT	TCTTTAATAT	TAACATCACC	4200
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TTCGTTTGAA	ACTAATTG	TGATTAGGTT	TGTTAAGTTT	TCTTTAACAT	CTGTGAAGCT	4380
TTCTTCTAAA	TATAATCTT	TGATTGCATT	AACTCTATAG	TCACCTAAC	GATTTAGATG	4440
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AGTATATTCT	TTTACATTAC	TTAAACCTTC	ACTGCTACAA	CACAAAGTTAA	AGTAATCTTT	4620
TGTACCGTCC	GCATAGTGAA	CAATAATT	ATTAGCTCA	TCTAGTTTG	TGATAAACTC	4680
ATTGTTGTT	ATCGCGTAA	CAGAAAGAAC	TTCTTAGTA	TTTAGATGGT	GTTCTTTATT	4740
TAATTATTA	CCTTGATATA	CAATATAATC	TTTATTGTA	AATGGTTA	ATTTTTCAAG	4800
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AGTAGTAACT	TTAGCATCAG	CTTGTCTTT	AGTTAATTAA	GTAAATGTTT	TAGATTCCCT	4980
AAATGATCTA	TTACCTGACG	AAATATCCCTC	TACCGCATAT	AAATCTTTA	TATGAGCACT	5040
AGCATAATCA	GAATCATCAA	CGTCGTTAGA	GCCGAATAAC	TCCTCTCCAC	GGATAATCTT	5100
AGCATAGCTG	ACAGAATTAC	TTACCGTACC	TACAGGCCAA	GTCTTACTTG	CTATTGCTCC	5160
AACTTCTACT	GGATTTGAAA	CATCTATT	ACCTTTACA	ACCGACTCAG	TTAGGAGAGC	5220
TTTTGTACCA	ATAAGATGGT	CTAGAGTTAA	TCCATAATCT	ACTTTAGGAA	CTAACAAAGCT	5280
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178

ATAGTTGTATCCTGCAATTCCCCTGTATGAGAGCCATTCCACTTGTAGAGTGTAGTTT	5400
GCCAAAGAAA GCAACATTTCATACCGAGT TCCATCATTCATATTATTTCAAAATCCAGC	5460
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TTTCATAGTA TTGGCTAATGATGCAATATTATCTTGACCAAAACGTTCTAATCTCTACATT	5580
TTCAAAATTCACATTATTTATCGTTGCGTTTGTTATCACAATGATGTTCCAA	5640
TTTCAGTAATA GCAAATTGTTTCCCTTCAGAACTTAAAGTTTCCTGTGATTCTTAGT	5700
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AACCGTAGTTCTCTATATTAAACGCTAGTAATGTTTTCTGATTTTGAAAGT	6000
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TCTTGGCGCGACTGGTCACCTGCTGTTCAACTTTGGTTCCTCTGTTGTTCTGTTG	7020
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179

TTCCCTCTACA GCCTTCTCTT CTTCAGGAGC TTCTGGTTGC TTTCTGGCT CGACTGGTGC	7200
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TACTTTAGGA AGGGTGTGCGT CAGTAGGTTT TACCTCCGAT TTTGGTTCTT CCTTTGGACT	7440
TTCTTCGTT TTAGGTGCTT CTTCTTTGG AGCTTCCCT GTCTCTACTA CTTGGTTTC	7500
TGTCTTAGCT TGCTCCTGAT TTGTTATTGA TTGAGGAGTC TCAACTTCGA CCACAGTCAC	7560
CTCTCCAGGT TTTGCTGAGG TTTCTTCTAA AACAGTGTCC AAGCCAAGCG TTTTGAGGAT	7620
GTCACCTGAT AGATAACCAA CATAGCGATA GCCCTCCATT TCAACAAACAC CCTCTCGACT	7680
AGCCAGCGCT AGGGTCGCAA CTGGGTCTAC AGCCCCCTGCA CTAGGAAGAA CTACCAATCC	7740
CATAGCTCCA ACTAGAAAGA CGCTAGCAAT TTTCTTCTC TTGTAGATTA AAAGCAAGCT	7800
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ACTAGCCAGT GCGTTACAT GGGCATGACC AATCTCTCTC AAAATAGGGC GAATCGGAAC	8220
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CTTGATAAAAT TCAACCTCAA CTGGATCCTG CATAAACTTA AAGGCTGCCA ACTGCCCGA	8340
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CTGACGTTCA ACAACGAGAG CCCGATTGAC ATGCTCACAA CCTTGAACGT CTAAATGGAT	8460
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GGCCCCCTGC TTCAAATTGG TCTTTCTAA AACATCTCC ACTACCTGAC GTGTTCTCT	8640
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CAGAATTGAC TGGGAGTTAG CTAGTTCTA TTCTATTTAT ATATATTTCA ACTTTCGTCC	8820
CTTTTTGGGG TCTAGAATCA ATCTTCATAT GGTAATTGGC TCCAAAATGA ACTTTGAGCC	8880

180

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TTTCATTAAT	TCGTATTCC	AGCTTATCTC	CATAGCGTTG	TTTCTGGATA	AAGAGATACT	9180
GGCGGACATG	ATTGATTTCG	TCAGAGAGAC	AAATCAAGTC	CTTGCCCTGA	TTGAGCGCCA	9240
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CAGCCATCCA	GATGATGGTG	TCCAAAGTGT	TATAGAGGAA	ATGTGGATTA	ATCTGGCTCG	9360
AAAGGGCTTG	AAGTTGGTAC	TGACGGGTG	TTTCTTCCTG	GCTACGAATA	GCTACCATCA	9420
ACTGATCAAT	CTGATCCAAC	ATAGCATTAA	ATTGGCGAGT	TACTTCTCTC	AGTTCATAGG	9480
CACCAACTTC	CTTGGCACGA	AGATTTGAG	CACCAAGAAC	ATTTCCAAC	ATGGTTCTC	9540
TCAAATCCTT	CAAAGGAGCA	ATCCAGCGTT	TAAGACTGAA	CCACACTAAG	CAGAGACAGA	9600
CAAGAAGAGA	TGTGACACTG	GCCCCAAGCA	AGGTCACAA	GAGCTGACTC	CGAACCTGGT	9660
CTAACTTTTC	CAATGATGAC	ACGCCAAGCA	CCGTCCAATC	AGTTCCCTGA	ATCTTCTCTT	9720
GACTGACGTA	GGATTGTA	CCAGGAGTAT	AACCCCTGACC	TGTATCGATG	TAGGTTTCA	9780
TAGCCTCCAT	TTTGCTAGAC	GAACTATAAA	CTGTGTGTTG	AGGATGGTAG	ACAAATTTCAT	9840
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CTTCCAGAGT	TTCATAAGAA	ATATCCAAAC	GAAGCACACC	AAGATTGGCT	CCCTTGCAT	9960
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AAACAGGCAT	AGCTCCCTGA	TGAATGGCCT	TTTGGTACCA	ATCCTCAGCC	ATCATATCAG	10080
AGGAAGTTTT	CATCTGCACA	CTGTCATCTG	TAGAAATGAC	CTGACCAGAT	TTGGTCACCA	10140
GCACAAACAGT	TTTCAAGTCC	TTATCTGACT	TCAAGATGGT	CAAAACAAA	TCTCGGATTTC	10200
CCTCGACCTT	GTCTTGACTG	GGATTCTCAG	CATAGGCCAG	AACATCCGTC	TGCTGGGTCA	10260
AACCAAGTCGA	GGTGGTTCT	AGTTTTTGAA	TATAAGACTG	AATAAAGTGG	CTAGTCTGGC	10320
TGATGGTCGT	TTGGCTGTTG	CCCTCAATGG	TGGCCTCAAT	GGCTGAAGAA	CTTGATTGAT	10380
AGTAGAAAGT	TCCAACCAGA	GCTAGGAGAA	TGAGAAAGAC	CAGAAAGATG	GAAATAACCA	10440
TTCTAACTAA	AAGAGAAGAA	CGCTTCATCG	GTCTTCTCCC	TTCTTAAACT	GACGAGGTGT	10500
CACACCTGCA	ATCTGCTTAA	AACGTTGGGT	AAAATAGTTC	ATATCTCAA	AACCAACCTT	10560
CTCTGCGATC	TCATAATCT	TCAGATCTGT	AGTTAAAAGC	AAGAGCTTGG	CTTGTAAAC	10620
ACGTTCTCTC	ACCAAGATAAT	CCTGAAAAGG	CAAGCCCAAC	TCTTTCTTAA	TCAAGGAAC	10680

181

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TTGTAACTGC TCTTCTTCT CTTCCCTGTC TAGTTTTGT TTGATTTCC CCAACATTTC	10860
CTCAATATCC TGACGAGAAA AGGGTTGAG CAGGTAGTCG TCCACACCTA GTTTGACAGC	10920
AGACAAGGCA TAATCAAAT CATCGTAACC TGTAAAG ACCAAATGAA CCTGAGGATA	10980
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CCAGTTGCTA TATCCACATA GATA CGGAT TCAAATTAT CCCAGTAACG GTTTGAGAAA	11820
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ACACCGACTAC GGTACTGGGT CCCCCACATCA TTTCCCTGTT TATTTTGCT GGTTGGATTG	12180
ATAATGCCGA AATAGTGAAG CAGGATTCC TTGAGAGAAA TTTGCTTGGC ATCATAGGTG	12240
ACATGGACGG TTTCTGCATG ACCTGTTGG TTAATCAATT CGTACTTGGT TGTGTTCTCCT	12300
CTACCATTTG CATAGCCTGA AACGGCATCC GTCACCCCGG GAACACGTGA GAAATATTCC	12360
TCCACTCCCC AGAACACAACC TCCAGCTAGA TAAATTCGT GCAAGTCTGC GTCTTTACTA	12420

182

ATTTCTGTTT	TTTCACTGC	TTTCCTCCT	TGGCTAACTG	CCGCCTTTTC	AATTTGCGAG	12480
GCATCTGTCT	GCCCTGCATT	TCGTATCAAT	AGAACATAGA	AACCGGTTAT	GGCTAGAAAA	12540
AATACTCCTA	GCAACAAGAA	GATTTTAAC	TTATCATTCA	TAAGACGCCT	CCTAGGCTAA	12600
TTCCCTCAAA	GTTTGCAAA	TTGCATCTTT	TTCCATGAAT	CCTGGATGTG	TTTTGACCAG	12660
CTTGCCTTCT	TTGTCTATAA	AGGCTTGGGT	TGGGTAAGAA	CGGACACCAT	AAGTTCCAA	12720
AAGTTTGCT	GATGGGTCAA	CTAGGACTGG	GAGTTTTA	TAATCCAATC	CCTTATACCA	12780
ATTCTTAAAG	TCCGCTTCAG	ATTGCTCTCC	CITATGTCCT	GGTGACACTA	CTGTCAAGAC	12840
CACATAGTCA	TCACCAAGCTT	CTTTAGCAAT	CTCATCCGTA	TCTGGAAAGAC	TAGCCAGACAA	12900
GATGGAACAC	CAAGAAGCCC	AGAATTGAG	ATAGACTTTC	TTGCCCTTGT	AATCAGATAA	12960
ACGGTAGGTC	TTGCCATCTA	CTCCCATCAA	TTCAAAATCA	GCCACCTCTT	TCCCTTAC	13020
TGCGCTTGT	TTACTAGCTG	TCTGCTCCGT	CTTCATTTCA	TCTTCGTTT	GGTGTTCACT	13080
AGTCACGGAC	TTGCCCTGAAC	AAGCCGTCAA	ACAAAGGAGC	GAACCTGCTC	CAAGAACACA	13140
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TTGAAGCATT	TCCAAACAGA	ACCAAGAACG	CCATCACAAAT	AATGAGAAAA	CCACCCACTT	13260
TTTGAGGAT	TCCCAGATAG	GGATGAAGTT	TTCGGAAATG	TTTCAAAACA	TAACTAGAGG	13320
TCAGAGCTAG	AAGCAAGAAT	GGTAGCGCCA	AGCCCAGCGT	ATACACCAAC	ATGAGACCAG	13380
CTCCCTGCCA	AGCTCCTGAA	CCACCTGAAG	CCGCCAAGGC	CAAAACAGAC	CCCAGAACCG	13440
GCCCCACGCA	AGGCCCTCAA	GCAAAACTAA	AGGTCAAGGCC	CAATAAAAAT	GCCTGACTAT	13500
AGCCCTTACC	ATTTGCCCC	TGTCCTTGCA	GTTGTAAGCCT	CTTTTCCCTTA	AAAGCCCT	13560
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ATTGGAACCA	AGAACATAA	AGCAAATCGC	CTAAAAAACCC	AGCTCCATAG	CCCAACAAAA	13680
TAAATATAAA	GGAAATTCCCT	GCTATAAAGG	CCAGAGTTCG	TAATAAACTA	GTAACGTGAGA	13740
TTGAAAATTT	GCGCTAGAA	GCCTGAGCAC	CATCCTTATC	ATCTAGTAAC	ACTCCTGTAT	13800
AGACCGGTA	CAAAGGTAAG	ATACAAGGAG	AAAAGAAGGA	TAGAATCCCT	GCCAAAAGA	13860
CACTTAGAAA	AAAGAAAATA	TGACCCATAA	AGTTCCCTCCT	ATCATTAT	TGATAGATT	13920
ATTATA						13926

(2) INFORMATION FOR SEQ ID NO: 6:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 20199 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:

CCCAGCAGAA	AAATGGCATT	TGGAGATAAT	GGAAATCGTA	AAAAAACTAT	GTGGAGAAA	60
ATAACCTTGT	TTATCGTGAT	TATCATGCTA	GTAGCAAGTT	TATTGGAAAT	TTTGCAACT	120
GCAATTGGTG	CCCTCAGTAA	TCTATAAAAT	AGATTCAAGA	AAATTTAGTG	ACTGGGATTT	180
CCCAGCCCTT	TTTTAAAGTG	AGAAGAAATA	ATGAGTATGT	TTTTAGATAC	AGCTAAGATT	240
AAGGTCAAGG	CTGGAATGG	TGGCGATGGT	ATGGTTGCCT	TTCGTCGTGA	AAAATATGTC	300
CCTAATGGAG	GCCCTTGGGG	TGGTGATGGT	GGTCGTGGAG	GCAATGTGGT	CTTCGTTGTA	360
GACGAAGGAC	TACGTACCTT	GATGGATTTTC	CGCTACAATC	GTCATTTCAA	GGCTGATTCT	420
GGTGAAAAAG	GGATGACCAA	AGGGATGCAT	GGTCGTGGTG	CTGAGGACCT	TAGAGTTCGA	480
GTACCCACAAG	GTACGACTGT	TCGTGATGCG	GAGACTGGCA	AGGTTTTAAC	AGATTTGATT	540
GAACATGGGC	AAGAATTAT	CGTTGCCAC	GGTGGTCGTG	GTGGACGTGG	AAATATTCTG	600
TTCGCACAC	CAAAATACTC	TGCACCGGAA	ATCTCTGAA	ATGGAGAACCC	AGGTCAGGAA	660
CGTGAGTTAC	AATTGGAAC	AAAAATCTTG	GCAGATGTGCG	GTGGAGTTAGG	ATTCCCATCT	720
GTAGGGAAAGT	CAACACTTT	AAGTGTATT	ACCTCAGCTA	AGCCTAAAAT	TGGTGCCTAC	780
CACTTTACCA	CTATTGTACC	AAATTAGGT	ATGGTTCGCA	CCCAATCAGG	TGAATCCTTT	840
GCAGTAGCCG	ACTTGCCAGG	TTTGATTGAA	GGGGCTAGTC	AAGGTGTTGG	TTTGGGAACT	900
CAGTTCTCC	GTCACATCGA	GCGTACACGT	GTATCCTTC	ACATCATTGA	TATGTCAGCT	960
AGCGAGGGCC	GTGATCCATA	TGAGGACTAC	CTAGCTATCA	ATAAAGAGCT	GGAGTCTTAC	1020
AATCTTCGCC	TCATGGAGCG	TCCACAGATT	ATTGTAGCTA	ATAAGATGGA	CATGCCTGAG	1080
AGTCAGGAAA	ATCTTGAAGA	CTTTAAGAAA	AAATTGGCTG	AAAATTATGA	TGAATTGAA	1140
GAGTTACAG	CTATCTTCCC	AAATTCTGGA	TTGACCAAGC	AAGGTCTGGC	AAACACTTTA	1200
GATGCTACAG	CTGAATTGTT	AGACAAGACA	CCAGAATTTC	TGCTCTACGA	CGAGTCCGAT	1260
ATGGAAGAAG	AAGCTTACTA	TGGATTGAC	GAAGAAGAAA	AAGCCTTGA	AATTAGTCGT	1320
GATGACGATG	CGACATGGGT	ACTTTCTGGT	AAAAAACTCA	TGAAACTCTT	TAATATGACC	1380
AACTTGATC	GTGATGAATC	TGTCTATGAA	TTTGCCCGTC	AGCTTCGTGG	TATGGGGTT	1440
GATGAAGCCC	TTCGTGCAGC	TGGAGCTAAA	GATGGGGATT	TGGTCCGCAT	TGGTAAATT	1500
GAGTTGAAT	TTGTAGACTA	GGAGACTGGT	ATGGGAGATA	AACCGATATC	TTTCCGAGAT	1560
CGGGATGGTA	ATTTGTTTC	CGCCGCAGAC	GTGGAGATG	AAAAGAAATT	GGAAGAACTA	1620

184	
TTTAATCGTC TCAATCCAAA TCGTGCCTTG AGATTGGCAC GAACTAAAAA GGAAAATCCA	1680
TCTCAGTAAA GAAGCTAAAA AATCCCGTGC CTCATCAGAC ACGGGATT TT GTGGTACGAC	1740
AGGCATGTAT AGCAAACGTGA ATCTGGAATA GCACAGCATA TCTTCTAAAA TATAGTAAAA	1800
TGAAATGAGA ACAGGACAAA TCGATCAGGA CAGTAAAATC GATTCTAAC AATGTTTAT	1860
AAGCAGAGAT GTACTATTCT AGTTCAATC AACTATATTG TTATAAATTG ATTTGAATT	1920
CAAAATTAAA TTGTTTGATT CTTATTCAA TTTGTTATAG TATATCTGAT GTCAAAGTTC	1980
TCGGCGAGTC AAATAGCGAT TCCCAAGCCT GACTATCGTG AGGTAGCGGA TTAAAATGGT	2040
CTGGGGATAG ACCGTTTAA GTCTGACGCT GGAAATAAGA ATTGTAGAA GAAGGGATAG	2100
CGAAATCGTG GCTCTACGAA CAGGAACGTG ATAATAAGGC GTATATAGCG GATAAGAGGG	2160
CATCAAACTC TAAAGTCCAA AAAGGTAGTC GTAACCTATA TGCGTAAATC ACGAGACTAA	2220
TTGAATTCTG ACTAAAGATTT TCTATTTCA CTGTAACCTT TTAACGCCCT TATATCTTGT	2280
ATACACGAGG AAAGATGTAC GACTTATCCC GTGAGGTCTA TCACTATAAA GAGAAAACGA	2340
CAGATAGAAG TGATCCTGAG TCACGGTTAT CTGTCGTGATA GGACGGTATG TATAAAACGC	2400
TTCTGTGAAC TGAGAGAAGG GGGAGAAGTT CTTGCTAAAA TTTAGTTGAA CAGCCGTATT	2460
CCGATACTTA GATAAGAGAT CTAGTCTTAG CTCCCTACTCA GTTTTAGGGG ATAAAAAAGG	2520
GGCAATAGCG ATTGAGAAA GATTATACTC TTGAAAATC TCTTCAAATC ACGTCAATAT	2580
CGCCTTGTGCG TATGTGTAGG ATACTGACTA CGTCAGTTC ATCTACAACC TCAAAACAGT	2640
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AATTCACTTA CTAACTCGTC AACTCTGATT TATCCAATAA AATTGAAAAG GATGGAAAAA	2760
AGGATAAATT TATGATATAC TTTATTTGA AGACCTTATT AGAAATCTG AAAGAGTATT	2820
GAAAACCTAG AATGAGAAA ATTGTTATCA ATGGTGGATT ACCACTGCAA GGTGAAATCA	2880
CTATTAGTGG TGCTAAAAAT AGTGTGTTG CCTTAATTCC AGCTATTATC TTGGCTGATG	2940
ATGTGGTGAC TTTGGATTGC GTTCCAGATA TTTGGATGT AGCCAGTCTT GTCGAAATCA	3000
TGGAATTGAT GGGAGCTACT GTTAAGCGTT ATGACGATGT ATTGGAGATT GACCCAAGAG	3060
GTTGTCAAA TATCCAATG CCTTATGGTA AAATTAACAG TCTTCGTGCA TCTTACTATT	3120
TTTATGGGAG CCTCTTAGGC CGTTTGGTG AAGCGACAGT TGGTCTACCG GGAGGATGTG	3180
ATCTTGGTCC TCGTCCGATT GACTTACACC TTAAGGCAGTT TGAAGCTATG GGTGCCACTG	3240
CTAGCTACGA GGGAGATAAC ATGAAGTTAT CTGCTAAAGA TACAGGACTT CATGGTGCAA	3300
GTATTTACAT GGATACGGTT AGTGTGGGAG CAACGATTAA TACGATGATT GCTGCGGTTA	3360
AAGCAAATGG TCGTACTATT ATTGAAAATG CAGCCCGTGA ACCTGAGATT ATTGATGTAG	3420

185

CTACTCTCTT	GAATAATATG	GGTGCCTATA	TCCGTGGGC	AGGAACTAAT	ATCATCATTA	3480
TTGATGGTGT	TGAAAGATTA	CATGGGACAC	GTCATCAGGT	GATTCCAGAC	CGCATTGAAG	3540
CTGGAACATA	TATATCTTTA	GCTGCTGCAG	TTGGTAAAGG	AATTCTGTATA	AATAATGTT	3600
TTTACGAACA	CCTGGAAGGG	TTTATTGCTA	AGTTGGAAGA	AATGGGAGTG	AGAATGACTG	3660
TATCTGAAGA	CAGCATTTC	GTCGAGGAAC	AGTCTAATT	GAAAGCAATC	AATATTAAGA	3720
CAGCTCCTTA	CCCAGGCTTT	GCAACTGATT	TGCAACAAACC	GCTTACCCCT	CTTTTACTAA	3780
GAGCGAATGG	TCGTGGTACA	ATTGTCGATA	CGATTTACGA	AAAACGTGTA	AATCATGTTT	3840
TTGAACTAGC	AAAGATGGAT	GCGGATATT	CGACAACAAA	TGGTCATATT	TTGTACACGG	3900
GTGGACGTGA	TTTACGTGGG	GCCAGTGTAA	AAGCGACCGA	CTTAAGAGCT	GGGGCTGCAC	3960
TAGTCATTGC	TGGGCTTATG	GCTGAAGGT	AAACTGAAAT	TACCAATATC	GAGTTTATCT	4020
TACGTGGTTA	TTCTGATATT	ATCGAAAAT	TACGTAATT	AGGAGCGGAT	ATTAGACTTG	4080
TTGAGGATTA	AACCGTAGAG	GTGTTTATGA	ATATTTGGAC	CAAATTAGCA	ATGTTTCTT	4140
TTTTGAAAC	GGATCGCTTG	TATTTGCGTC	CTTTCTTTT	TAGTGTAGT	CAGGACTTCC	4200
GCGAGATAGC	TTCAAATCCA	AAAAATCTTC	AATTTATTTT	CCCAACGCAG	GCAAGTCTGG	4260
AAGAAAGTCA	ATATGCACTG	GCCAATTACT	TTATGAAGTC	CCCTTGGGA	GTGTGGCAA	4320
TTTGTACCCA	AAAAATCAA	CAAATGATTG	GTTCTATTAA	ATTTGAGAAG	TTAGATGAAA	4380
TCAAAAAAGA	AGCTGAGCTT	GGCTATTTTT	TGAGAAAAGA	TCCTTGGTGC	CACCAATTAA	4440
TGACAGAGGT	TGTTAGAAA	ATTGTCAGC	TTCTTTGGA	GGAAATTGGC	TTAAAACAAT	4500
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TTGGAGGAAT	TACCACTAGG	AAAAAGGGTC	AGTGTTCGTA	GCATTTGAA	TCATCTAGGA	4740
GTTAGTGATG	GAACAGCCTA	TCGGGCTATT	AAAGAAGCTG	AAAACCGTGG	AATTGTGGAG	4800
ACCCGTCTA	GAAGTGGAAC	AATTCTGTGT	AAATCCCAGA	AGTTGCTAT	AGAGAGATTA	4860
ACGTTGCTG	AAATTGCAGA	AGTGAATTCT	TCTGAGGTT	TGGCTGGCA	AGAAGGTTA	4920
GAGAGAGAAT	TTAGTAAGTT	TTCAATTGGT	GCCATGACTG	AACAAAATAT	CTTGTCTTAC	4980
CTTCATGATG	GGGGCTCTT	GATTGTCGGA	GACCGAACCC	GTATTCTAGTT	GCTAGCCTTG	5040
GAAAATGAAA	ATGCACTTCT	GGTTACAGGG	GGATTCAGG	TTCATGATGA	TGTGCTTAA	5100
CTGGCCAATC	AAAAAGGGAT	TCCCTGTTCTA	AGAAGTAAGC	ATGATAACCTT	TACCGTCGCG	5160

186

ACCATGATCA ATAAAGCCTT	GTCAAATGTC CAAATCAAGA	CTGATATTCT	CACAGTTGAG	5220		
AAACTTATC	GCCCTAGTCA	TGAGTATGGT	TTTCTGAGAG	AGACAGATAC	AGTTAAAGAT	5280
TATTTGGACT	TGGTCGTAAC	GAATCGTAGC	AGCCGTTTCC	CTGTTATCAA	TCAACATCAG	5340
GTCGTTGTTG	GTGTTGTAAC	CATGAGAGAC	GCTGGTGATA	AATCACCAAG	CACGACAATT	5400
GATAAGGTTA	TGTCTCGTAG	TCTATTTTG	GTGGATTAT	CGACAAATAT	TGCCAATGTG	5460
AGTCAACGGA	TGATCGCAGA	AGACTTGAA	ATGGTACCAAG	TTGTTGAAAG	CAATCAAAC	5520
TTGCTTGGCG	TTGTGACGCG	ACGAGATGTC	ATGGAGAAGA	TGAGCCGTT	CCAAGTTTCG	5580
GCTCTACCAA	CTTTTCTGA	GCAGATTGGA	AAAAAGCTCT	CTTATCACCA	TGATGAAGTA	5640
GTCATTACAG	TGGAACCCCTT	TATGCTAGAA	AAAAATGGAG	TTTGGCTAA	TGGTGTATTG	5700
GCAGAAATTC	TGACCCACAT	GACCCGATT	AGTTGTTAAT	AGTGGTCGCA	ATCTCATTAT	5760
CGAGCGAGATG	CTGATCTACT	TTTGCAGGC	TGTTCAGATA	GATGATATAT	TGCGCATTCA	5820
GGCACGGATT	ATTCACTATA	CGAGACGGTC	AGCTATAATT	GATTACGATA	TTTATCATGG	5880
TCACCAAGATT	GTTCAAAAG	CAAATGTGAC	TGTTAAATT	AATTAGAAC	TAGGAGAAAA	5940
GATGATAACA	TTAAATCAG	CTCGTAAAT	CGAAGCTATG	GACAAGGCTG	GTGATTTCT	6000
AGCAAGTATT	CATATAGGCT	TACGTGATT	GATTAAGCCA	GGCGTAGATA	TGTGGAAAGT	6060
TGAAGAATAT	GTCCGCCGTC	GTTGTAAGA	AGAAAATTTC	CTTCCACTTC	AGATTGGGTT	6120
TGACGGTGCC	ATGATGGACT	ATCCTTATGC	TACCTGTTGC	TCTCTTAACG	ATGAAGTGGC	6180
TCACGCTTTC	CCTCGTCATT	ATATCTTGAA	AGATGGTGAT	TTGCTCAAAG	TTGATATGGT	6240
TTTGGGAGGT	CCCATTGCTA	AATCTGACCT	AAATGTCCTCA	AAATTAAACT	TCAACAAATGT	6300
TGAACAAATG	AAAAAATACA	CTCAGAGCTA	TTCTGGTGGT	TTAGCAGACT	CATGTTGGCC	6360
TTATGCTGTT	GGTACACCGT	CCGAAGAAGT	AAAAAACTTG	ATGGATGTAA	CCAAAGAAC	6420
TATGTACAAG	GGTATTGAGC	AAGCTGTTGT	TGAAATCGT	ATCGGTGATA	TCGGTGCAGC	6480
TATTCAAGAA	TACGCTGAAA	GTCGTGGTTA	CGGTGTAGTG	CGTGATTG	TTGGTCATGG	6540
TGTTGGCCCA	ACTATGCAAG	AAAAGCAAT	GGTCTTAAC	TATGGTATTG	CAGGTCGTGG	6600
ACTCCGTCTT	CGTGAAGGAA	TGGCTTAAAC	CATTGAACCA	ATGATCAATA	CAGGCATTG	6660
GGAAATTGAT	ACAGATATGA	AAACTGGTGTG	GGCGCATAAG	ACCATTGACG	GTGGATTGTC	6720
ATGTCAGTAT	GAACACCAAT	TTGTCATTAC	GAAAGATGGA	CCTGTTATCT	TGACTAGCCA	6780
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AAAATCCAGT	AGATCTTTC	ATAATAAAAC	GCATTGTATC	AAGTGTAGG	GGCTGATATC	6900
ATGCGTTTT	CTGCTTTAA	GATTTTTCC	AACTCTGTTT	GTAAGCGCAT	CATAACAAAG	6960

GGTCTAGGAT	TCAGGGCTCT	CCTCCTATAT	ACTATTAGTA	AAGTAAAAC	AAGGGAGGAT	7020
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AAAAAGAGAA	ATGGAATTCT	GTTTAATGTT	CGTATTGCCA	ATCTTAAATT	TACTTTTTA	7140
TATTATACTT	CCTGCCAAC	AAAATATGGT	ATAGTAGTTC	TATGAATGAT	GAAGCAAGTA	7200
AACAACAAAC	TGATGCCAGA	TTTAAGCGTC	TTGTTGGTGT	TCAGCGTACC	ACTTTGAAAG	7260
AGATGTTAGC	TGTATTAAAA	ACAGCTTATC	AACTTAAACA	CGCAAAAGGT	GGACGAAAAC	7320
CTAAATTAAAG	CCTAGAAAGAC	CTTCTTATGC	CCACTCTTCA	ATAGTGCAG	AATATCGAAC	7380
TTATGAAAGAA	ATTGCGGCTG	ATTTGGTAT	TCACGAAAGC	ACCTTTATCC	GTCGGAGCCA	7440
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TGAGGACACG	GTAATGATTG	ATGCCACGGA	AGTAAAATC	AATGCCCTA	AAAAAACAAAT	7560
TAGCGAATGA	TTCTGGTAA	AAGAAATTTC	ACGCTATGAA	GGCTCAAGCG	ATTGTCACAA	7620
GTCAAGGGAG	AATTGTTCT	TTGGATATCG	CTGTGAACTA	TAGTCATGAT	ATGAAGTTGT	7680
TCAAAATGAG	TCGCTAGAAAT	ATCGAACAAAG	CTGGTAAAT	CTTGGCTGAC	AGTGGTTATC	7740
AAGGGCTCAT	GAAGATATAT	CCTCAAGCAC	AAACTCCACG	AAATCCAGC	AAACTCAAGC	7800
CGCTAACACG	TGAAGATAAA	GCCTATAACC	ATGCGCTATC	TAAGGAAAGA	AGCAAGGTTG	7860
AGAACATCTT	TGCCAAAGTA	AAAACCTTA	AAATATTTTC	ACAAACCTAT	CGAAATCATC	7920
GTAAACGCTT	CGGATTACGA	ATGAATTGA	GTGCTGGTAT	TATCAATCAT	GAACTAGGAT	7980
TCTAGTTTG	CAGGAAGTCT	ATTGAGGTAT	TGAGCTAGTT	TATGAAAAAA	TTGGGTGAAA	8040
AGTCGAGTGT	TTTAGAAACC	CACAGTGTAG	TATTCTAGTT	TCAATCCACT	ATATTTGCT	8100
ACTCCCCGTA	AAGTTCTAT	TTTCCCTGAT	TTCGTGATATA	ATAGAAATAT	TGACTTCAAG	8160
AGTAAGGAAG	AGAAGATGAA	CGCATTATTA	AATGGAATGA	ATGACCGTCA	GGCTGAGGCC	8220
GTGCAAACGA	CAGAAGGTCC	CTTGCTAATC	ATGGCAGGGG	CTGGTTCTGG	AAAGACTCGT	8280
GTMTTGACCC	ACCGTATCGC	TTATTTGATT	GATGAAAGC	TGGTCAATCC	TTGGAATATC	8340
TTGGCCATTA	CCTTACCAA	CAAGGCTGCG	CGTGAGATGA	AAGAGCGTGC	TTATAGCCTC	8400
AATCCAGCGA	CTCAGGACTG	TCTGATTGCG	ACCTTCCACT	CCATGTGTGT	GCGTATTTG	8460
CGTCGCGATG	CGGACCATAT	TGGCTACAAT	CGTAATTAA	CAATTGTGGA	TCCTGGTGAA	8520
CAGCGAACGC	TCATGAAACG	TATTCTCAA	CAGTTGAAC	TGGACCCCAA	AAAATGGAAT	8580
GAACGAACTA	TTTTGGGAC	CATTCCAAT	GCTAAGAATG	ATTTGATTGA	TGATGTTGCT	8640
TATGCTGCC	AAGCTGGCGA	TATGTATAAG	CAAATTGTGG	CCCAGTGTAA	TACAGCCTAT	8700

188	
CAAAAAGAAC TTTCGTCAGTC TGAATCCGTT GACTTTGATG ATTTGATTAT GCTGACCTTG	8760
CGTCTCTTTG ATCAAAATCC TGATGTTTG ACCTACTACC AGCAAAAATT CCAATACATC	8820
CACGTTGATG AGTACCAAGA TACCAACCAC GCTCAGTACC AATTGGTCAA ACTCTTGGCT	8880
TCCCCGTTTA AAAATATCTG TGTGGTTGGG GATGCGGACC AGTCTATCTA CGGTTGGCT	8940
GGTGCTGATA TGCAGAATAT CTTGGACTTT GAAAAGGATT ACCCCAAAGC CAAGGTTGTT	9000
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AAAAATAATA AAAATCGCG TCCTAAAAAT CTCTGGACTC AAAACGCTGA TGGGGAGCAA	9120
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GATGAACCTTA GTCGCAGTCA AAACCTTCCTT CATAAGGATT TTGCAGTTCT CTATCGGACT	9240
AATGCCCAAGT CCCGTACAAT TGAGGAAGCC CTGCTCAAGT CTAACATTCC TTATACCATG	9300
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CTTATTGCTA ATTTGAGTGA CAATATTAGT TTTGAGCGTA TTATCAACGA GCCTAAACGT	9420
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ATCTGGGATT TTGCCAATAT GATGCTTGAT TTGCGGGAGC AGCTAGACCA CTTAACGATT	9600
ACAGAGTTGG TTGAGTCCGT CCTAGAAAAA ACAGGTTATG TCGATATTCT TAACTCCAA	9660
GCGACTCTAG AAAGCAAGGC ACGGGTTGAA AATATCGAAG AGTTTCTTTC TGTTACGAAG	9720
AACTTGATG ACACCACGGA TGTGACAGAA GAGGAAACTG GTCTGGACAA ACTGAGTCGT	9780
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GAAGAGCGCC GTCTAGCCTA TGTAGGTATC ACGGCGTGCAG AGAAAATTCT CTATCTGACC	10020
AATGCCAACT CACGCTTGCT TTTTGGTCGT ACCAATTATA ACCGTCCGAC TCGTTTTATT	10080
AACGAAATCA GTTCAGACTT GCTTGAGTAT CAAGGTCTGG CTCGTCCTGC AAATACAAGC	10140
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GCTCTTCAAG ACCGTAAACG CGGTGCTGCC CCAAAATCAA TCCAGTCAAG CGGTCTTCCA	10260
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GCTAGGCAGG AATTGAAAAT CAATTCCCA GAAGTAGGTT TGAAAAAAACT TTTAGCCAGT	10440
GTGGCTCCAA TTGAGAAAAA AATCTAATTT TCCATCCCTC TCACGAATAA TAAAGTGAGG	10500

AGGATTTTA TGTACAGTAT TTCATTCAA GAAGATTAC TATTACCAAG AGAAAGGCTG	10560
GCCAAGGAAG GAGTTGAAGC GCTTAGTAAC CAAGAGTTGC TAGCTATTTT ACTCAGGACA	10620
CGAACACGTC AAGCTAGCGT TTTGAAATT GCCCAAAAG TCTTGAACAA TCTTTCAAGC	10680
CTAACGGATT TGAAAAAAAT GACCTGCAG GAATTGCAGA GTTTGTCTGG TATTGGCGT	10740
GTAAAGGCC TAGAATTACA AGCTATGATT GAACCTGGGC ATCGTATTCA CAAACACGAC	10800
ACTCTGAAA TGGAAAGTAT TCTCAGCAGT CAAAGTTGG CCAAGAAGAT GCAGCAGGAA	10860
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ATTCTTCACT ATGCAATCAA GCATATGGCG ACTTCTCTTA TCTTGGTCCA CAATCATCCT	11040
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TTATCTTGG GACGATTTTC AAAAAGAAGT TCTGGATGCC ATTGGACACC GAGAAAGGCC	11280
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TCAACAGAAG AATCTGCCA ATGGTCTTCG ATATCTTGGT ACAAAAGTTCC ACCCATGGCA	11520
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ATCCCTCCAG CATCTTAAAC GCCTTCAACA AAGCCTTTG CTGCGTAGCT CATCATGATG	11820
TCATCATCTG GATGAGTTT TTCGTTTCCT GTAACTCCAA TAACTGGTTT TTTCATAAAA	11880
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TCAAATCGAC ATACTGAACG ACCACGTCTT TTGGTAAATG CAGATGGACT GGTGAAAAAC	12000
TGAGAATTCC TTTCACACCA GCATCAACCA AGAGATTAGC AACCTTGTGT GACTTGACCGC	12060
TGGGAACAGT TAGGATAGCA GTCTTCACAT CAGCATCCTT GATTTTATCC TTGATCTGAG	12120
AAATCCCGTA AATGGGAATC CCGTCAGGAG TTTGGGTACC GACTTCAGGA TGGTCGTCTA	12180
GGTCAAAGGC CATGATAATC TTCATCTTGT TACGTTCGTG GAAGCGGTAG TGGAGAAGGG	12240

190

CATGGCCCAT	ATTTCCAATA	CCAACCAGCA	TGACATTGGT	AATAGAGTTG	TCATTGAGCA	12300	
AATCGGC	AAA	AAATGTCATT	AGTTTTTGA	CATCATAGCC	AAAACCACGA	CGACCAAGTT	12360
CACCAAAATA	GGAAAATCA	CGACGTACGG	TCGCTGAATC	AATACCGATA	GCCTCTGCAA	12420	
TTTGCTTAGA	GTTGGCACGT	TCAATCTTT	CTGCATGAAA	TCTCTTAAAAA	ATTGATAGT	12480	
AGAGAGAGAG	TCTTTTGCT	GTAGCTTTG	GAATAGCAA	CTGTTATCT	TTCACAAAAT	12540	
CACAACCTT	CTATTCTCT	ATTTTATAGA	AACATTGTGA	AAAATCAAC	AAAATAAGA	12600	
AAAAACTAAG	AAAATCTTA	GTTTGATGT	AAAAAATCTG	CATGAGATAG	AAAACGGTAG	12660	
AGGTCTCCGA	CCAGCCCCCTG	ATAAACTTTT	TTGCCCTAA	AAGTCAGAGA	AGTCACATAA	12720	
AGTGTATCTG	GTAAGGTTAC	ACATCCTGAC	AAAGTCAACA	TGAGAGCCTC	ATGATCCTCA	12780	
TACTTGAGAG	TACGCTCTAC	ATGATAGCAG	TCCTTATAGG	TCAGTTCAA	CATTMTGGCT	12840	
CTATCTTCC	GATTTGTAA	AGACACCACG	TTCTACCAAG	CTATCCATGA	GGAAGTAGAA	12900	
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GAAGCCTTCA	CCGCTGTTG	GCACCTTTTC	CAAAGGCAG	GTCAGTTCAT	AGTTACCAAC	13080	
CTTAGTTCA	AAAAGGTGT	TATCTTGAG	GGTGAATT	TTAACAGAAG	GGCTAAGAGT	13140	
GTAATCGTAA	CGACAA	TTT	TTAACTGAAT	GATTTTTCA	AATGCCATAT	GGCTAACCTC	13200
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ACAAACTGAT	GCGAAGGGAT	TCCTTCAAGC	GTTCTGAATT	TGCGCCATAC	ATGGCTTCAA	13320	
GAACATGGCT	GGATGGACA	ACGCC	TACAGGCTGA	GCCAGTAGAG	ATTGAAATTC	13380	
CAGCTAAATC	TAGCCGAAGG	AGTAAGAGGT	CATT	TTCTGAG	ACCAGGAAT	CCAATATTGA	13440
GAACATAAGG	GAGATGATGT	TTTCTCTAT	TCAGGTAA	CTGAATGCC	TCCAGCTCTG	13500	
CCAGAAAGGC	AGTTTCTAGA	TTTGTACAT	CTTGAAT	TTCTTCTTGT	TTTCTAGGT	13560	
CTTCTTTAG	GGCTGCAACC	ATGCC	TACAA	TGGCAGGAG	ATTTTCAGTT	CCTGCACGTT	13620
TTTTCTGTT	CTGGCTCCG	CCATGTAGAT	AGGAATCAA	GTCCATGCTA	GATGCGTAGA	13680	
GAAAACCGAT	TCCCTTAGGA	CCATGGAATT	TGTGGCAGA	ACCAAGTGAGA	AAATCAATGC	13740	
CCAATTCTTC	TGAATGAATT	GGGATTTAC	CAATAGCCTG	AACTGCATCA	ACATGATAGG	13800	
CAGCAGGGTG	TTGCTTGAGT	ATTTGGCAA	TTTCAGCGAT	GGGCAGTAGG	TTTCTGTCT	13860	
CATTATTGAC	AAACATGGTA	GAACCAAA	TCGTATCGTC	ACGTAAGCC	TTTGAAATT	13920	
GCTGGGCTGT	GATTCTTGA	TTTCTGGCT	GGATAATGGT	TGCTTCAAAC	CCAAAGTGTT	13980	
GAACCAAGTA	ATCAATTGTT	TCAAGGACAG	CATGGTGCTC	GATGGCAGTT	GTGATGATAT	14040	

191

GTTCCTCCTTG	TTCTTGGTGA	CGAACAGT	AGCCAATGAT	GGTAGTATTA	TTGCCTTCAG	14100
TCCCACCCAGA	AGTAAAAAAG	ATATGTTGAG	GTTCCTGCCT	TAGTAACGG	GCTAGTTCC	14160
GACGGGCTTC	TCGCAAGAGT	TTGCCAGCTT	GACGACCATG	ACCATGAATA	CTAGAAGGAT	14220
TTCCGTGGGT	TTCTTGCATA	ACCTTGGTCA	TAGCTGAAAT	AGCAACTGCT	GACATAGGAG	14280
TCGTTGCAGC	ATTGTCCAAA	TAAATCAAAG	AATCACCTTA	TTTCTTTTA	TTGTAGGCAA	14340
AGAGTGGGCT	GACTGGTTT	CTTTCGTGAA	TACGGACGAT	AGCATCACCA	ATTAACACTCAC	14400
TAGCAGTGAT	GTAGCATAACA	TTTTTAGGAG	TTTTTCTTT	TGTTGCTACT	GAATCAGTCA	14460
CAAGAAATTTC	TTTAATATTA	GTATTGTCAA	GAAGCTCAGC	AGCTCCCTCG	ACGAAGAGAC	14520
CGTGGCTAGA	AACAGCATAA	ATTCTGTAG	CTCCTTCACG	TTCAACGATT	TTAGAAGCTT	14580
CAGAGAAGGT	ACGTCCTGTA	TTTAAATAT	CATCAATCAA	GATAGCTTTC	TTACCTTCA	14640
CATCACCAAT	AATATAACCT	TCGTTACGAG	TTGCATCGTC	TTGAGGGTAG	TCGATAATGG	14700
CGATAGGAGC	ATCAAGATAT	TCAGCCAGGC	TACGGCGACG	TTTGACACCT	GAATTTTTAG	14760
GGCTAACGAC	AACAACATCT	GAACCAAGCA	ATCCCTTATC	GCAGTAATGT	TTTGCGAATA	14820
GGGGAAACGT	GAAAAGATTA	TCCACTGGAA	TATCAAAGAA	ACCTTGAACC	TGAACGGCAT	14880
GCAAATCAAG	AGTCAGGATA	CGATCAACTC	CAGCCTTAAC	CAGCATATTG	GCAACTAGTT	14940
TTGCTGTAAG	TGGCTCACGA	GGACAAAGCA	TGCGGTCTTG	ACGTGCATAG	CCAAAATATG	15000
GAAGGACAAC	GTTGATACTG	TGGGCACCTTG	CACGCACACA	AGCATCGACC	ATGATTAACA	15060
ATTCCATTAG	GTGGTTGTTG	ACAGGGAAAC	TTGTTGATTG	GATGATGTA	ACATCATAAC	15120
CACGGACACT	TTCTTCGATA	TTTACTTGGA	TTTCTCCGTC	TGAAAATTGA	CGTGATGATA	15180
GTTCCTCAAG	TGGGACACCA	ACAGCTTGGG	CAATTTTTG	TGCAATCTCT	TGGTTAGAGT	15240
TGAGTGGAA	AAGTTTCATG	TTTTTTCTAT	CTGACATTAT	AGACCGCTCT	CTGTAAACTT	15300
TATAAACCT	AGTTTATTT	ACCTTACATA	TATGAACCTGG	GATTGTTGTA	TTTTTATCTT	15360
TTCTATTTA	CCAAAAATG	GAGATTATTT	CAGCTATTT	TCATACTTT	GACAAATCGA	15420
ACCAATTTCG	AAGGAGCTTT	TTGATAGGAA	ATCTGATTT	TCTCTAAAAA	TTGTCGAAAA	15480
TCCTGTTGC	CTTGCTCATG	ATTTTCCACT	TCAAGCTCCA	ATTCGTAATC	TGTTATATCA	15540
AAGTATCGGC	TCTGATCCAG	TGCCATGAGA	CCAATAGCTG	TTTCATTTTC	ATAGCGAAGC	15600
GTTGTTAGAC	AACCAAGAAC	CTGCCAGTTC	TTACTTTGGA	TACCATGTTT	CGCCAATTCA	15660
TCCAGTACTA	GCCCTTGAGG	AACTCTTCC	TTACTCAGAT	AGTTCTCAGC	ATCTTTAGT	15720
TGCAATTTC	GGTTGTATTTC	CATGTTTCCA	ACACTCTGCG	GGACTTTGAG	TGTCAACTCA	15780

192	
GCCCAGTCTT CAAAGGTTCG AATGCCATA GCGACTTCT TTTCTCGAG TTCAAAATCA	15840
GGCGTGTGCA TGTAGTAATT TGTGTTGAAGA ACAGGGAGTGA CACCTGTGAA CTGGTCTTT	15900
AGACGATTGT ATTCAATCTT TTTCAATAGT GTTTCAATT CAATTTCTAA ATGTTTCATT	15960
TTTCTTACCT TTTTTTATCG TTGAAAGCGG ATTTATGGTA TAATAAGCAT TGTATTTATT	16020
GTATATGAAT CTGGAGAAAA AATCAAAGAT ATTTTGACG GATAATATGA GAACAAGGGA	16080
GAATATATGA CCTTAGAATG GGAAGAATTT CTAGATCCTT ACATTCAAGC TGTTGGTGAG	16140
TTAAAGATTA AACTCGTGG TATTCGTAAG CAATATCGTA AGCAAAATAA GCATTCTCCA	16200
ATTGAGTTTG TGACCGGTG AGTCAAGCCA ATTGAGAGCA TCAAAGAAAA AATGGCTCGT	16260
CGTGGCATTA CTTATGCGAC CTTGGAACAC GATTTGCAGG ATATTGCTGG CTTACGTGTG	16320
ATGGTTCACT TTGAGATGA CGTCAAGGAA GTAGTGGATA TTTTGACAA CGGTCAAGGAT	16380
ATGCCAATCA TACAGGAGCG AGATTACATT ACTCATAGAA AAGCATCAGG CTATCGTCC	16440
TATCATGTGG TAGTGAATA TACGGTTGAT ACCATCAATG GAGCTAAGAC TATTTGGCA	16500
GAAATTCAA TTCGTACTTT GGCCATGAAT TTCTGGCAA CGATAGAACAA TTCTCTCAAC	16560
TACAAGTACC AAGGGGATTT CCCAGATGAG ATTAAGAACG GACTGGAAAT TACAGCTAGA	16620
ATCGCCCCATC AGTTGGATGA AGAAATGGGT GAAATTGCGT ATGATATCCA AGAAGCCCAG	16680
GCACATTGGATGAG TAGAAAATTA AATGACGGTG TAGGAAACAG TGACGATACA	16740
GATGAAGAAT ACAGGTAAAC GAATTGATCT GATGCCAAT AGAAAACCGC AGAGTCAAAG	16800
GGTTTTGTAT GAATTGCGAG ATCCGGTAA GAGAAATCAG TTTATACTCA ATGATACCAA	16860
TCCGGATATT GTCATTTCCA TTGGCGGGGA TGGTATGCTC TTGTCGGCCT TTCATAAGTA	16920
CGAAAATCAG CTTGACAAGG TCCGCTTTAT CGGTCTTCAT ACTGGACATT TGGGCTTCTA	16980
TACAGATTAT CGTGATTTG AGTTGGACAA GCTAGTGACT AATTTGCAGC TAGATACTGG	17040
GGCAAGGGTT TCTTACCCCTG TTCTGAATGT GAAGGTCTTT CTTGAAAATG GTGAAGTTAA	17100
GATTTTCAGA GCACTCAACG AAGCCAGCAT CCGCAGGTCT GATCGAACCA TGGTGGCAGA	17160
TATTGTAATA AATGGTGTTC CCTTTGAACG TTTTCGTGGA GACGGGCTAA CAGTTTCGAC	17220
ACCGACTGGT AGTACTGCCT ATAACAAGTC TCTTGGCGGT GCTGTTTAC ACCCTACCAT	17280
TGAAGCTTTG CAATTAACGG AAATTGCCAG CCTTAATAAT CGTGTCTATC GAACACTGGG	17340
CTCTTCCATT ATTGTGCCATA AGAAGGATAA GATTGAACCTT ATTCCAACAA GAAACGATTA	17400
TCATACTATT TCGGGTGACA ATAGCGTTTA TTCTTTCCGT AATATTGAGC GTATTGAGTA	17460
TCAAATCGAC CATCATAAGA TTCACTTTGT CGCGACTCCT AGCCATACCA GTTTCTGGAA	17520
CCGTGTTAAG GACGCCCTTA TCGGCGAGGT GGATGAATGA GGTTTGAATT TATCGCAGAT	17580

193

GAACATGTCA AGGTTAAGAC CTTCTTAAAA AAGCACGAGG TTTCTAAGGG ATTGCTGGCC	17640
AAGATTAAGT TTGAGGTGG AGCTATTCTG GTCAATAATC AACCGCAAA TGCAACGTAT	17700
CTATTGGACG TTGGAGACTA CGTTACCAATT GACATTCCCG CTGAGAAAGG CTTTGAAACC	17760
TTGGAGGCTA TTGAGCTTCC ATTAGATATT CTCTATGAGG ATGACCACCT TCTAGTCTTG	17820
AATAAACCT ATGGAGTGGC TTCTATTCTC AGTGTCAATC ACTCTAATAC CATTGCCAAT	17880
TTTATCAAGG GTTACTATGT CAAGCAAAAT TATGAAAATC AGCAGGTTCA CATTGTTACC	17940
AGACTAGATA GGGATACTTC TGGCTTGATG CTCTTGCCA AGCACGGTTA TGCCCATGCA	18000
CGATTAGACA AGCAGTTGCA GAAGAAATCT ATCGAGAAC GCTACTTTGC TTTGGTTAAG	18060
GGAGATGGAC ATTTGGAGCC AGAAGGGAA ATTATTGCTC CGATTGCGCG TGATGAAGAT	18120
TCCATTATTA CCAGACGAGT GGCTAAAGGC GGAAAGTATG CCCATACTTC ATACAAGATT	18180
GTAGCTCTT ATGGAAATAT TCACCTGGTC TATATTCAAC TGCACACTGG TCGAACCCAT	18240
CAAATCCGAG TCCATTTC TCATATCGGT TTCCCTTGC TGGGAGATGA TTTGTATGGT	18300
GGTAGCTGG AAGATGGTAT TCAACGTCAG GCTCTGCATT GCCATTACCT ATCCCTTTAT	18360
CATCCATTTC TAGAGCAAGA CTTGCAGTTA GAAAGTCCCT TGCCGGATGA TTTTAGTAAC	18420
CTTATTACCC AGTTATCAAC TAATACTCTA TAAAAACTGT CTCAGAGTAT AATTATTATC	18480
TTAAAGGAGA AAACTCATGG AAGTTTTGA AAGTCTCAA GCCAACCTTG TTGGTAAAAA	18540
TGCTCGTATC GTTCTCCCTG AAGGGGAAGA GCCTCGTATT CTTCAAGCAA CAARACGCTT	18600
AGTAAAAGAA ACAGAAGTGA TTCTGTTT GCTGGAAAT CCTGAAAAAA TTAAAATTAA	18660
TCTTGAAATT GAAGGAATCA TGGATGGTTA TGAGGTCACT GACCCTCAAC ATTATCCTCA	18720
ATTTGAAGAA ATGGTTCTG CCTTGGTGG ACGTCGCAAG GGCAAAATGA CTGAAGAAGA	18780
TGTACGCAAG GTTTGGTTG AAGATGTCAA CTACTTTGGT GTGATGTTGG TTTACTTGGG	18840
CTTGGTTGAT GGAATGGTGT CAGGAGCGAT TCACTCAACA GCTTCAACAG TTCGCCAGC	18900
TCTACAAATC ATCAAAACTC GTCCAATGT AACTCGTACT TCAGGAGCCT TCCTCATGGT	18960
TCGTGGTACG GAACGTTACC TATTTGGAGA CTGTCGCCATT AACATCAATC CAGATGCAGA	19020
AGCCTTGGCT GAAATTGCCA TCAACTCAGC AACACAGCT AAGATGTTG GCATCGAAC	19080
TAAAATTGCC ATGTTGAGCT ATTCTACTAA AGGTTCAGGG TTGGTGGAAA GCGTTGATAA	19140
GGTCGTTGAA GCAACTAAA TTGCTCACGA CTTGCGCCT GACCTTGAAA TCGATGGTGA	19200
GTGCAATTGATGTCAGCT TTGTTCTGA AACTGCAGCT CTGAAAGCTC CTGGAAGTAC	19260
GGTAGCTGGT CAAGCAAAATG TCTTCATCTT CCCAGGTATC GAGGCAGGAA ATATTGGTTA	19320

194

CAAGATGGCT	GAACGCCCTGG	GTGGCTTGC	GGCTGTAGGA	CCTGTTTGC	AAGGTTAAA	19380
CAAGCCAGTT	AATGATCTT	CTCGTGGATG	TAATGCAGAT	GATGTTACA	AGTTGACCT	19440
CATCACAGCA	GCTCAAGCAG	TTCATCAATA	GTGAAAAC	TAAAGTGATA	TACTATGCTA	19500
TACTGTAGTT	ATGAAACTAT	GTACGAAAAG	CACTGCCATT	AATTCCCTGAG	AACTAAATT	19560
CTGATGGTG	TCAAAAAGGA	AAACTTCAA	GGCATGATAT	CCTGTCTATA	CACGACCTAT	19620
AGAAAATCTGT	AATATACATA	TCCGTAAAAC	GATAAAATTCC	CTTTTGATT	TTAAATGAGT	19680
ATGAAAAGAG	AATTTTTTGG	CTCTTTGTCA	ACTGTAGTGG	GTTGAAGAAA	AGCTAAGCTC	19740
GAGAAAGGAC	AAATTTCATC	CTTCTTTTT	TGATATTCA	AGCGATAAAA	ATCCGTTTT	19800
TGAAGTTTC	AAAGTTCCGA	AAACCAAAGG	CATTGCGCTT	GATAAGTTG	ATGAGATTAT	19860
TGGTCGCTTC	CAGTTGGCG	TTAGAATAGT	GTAGTTGAAG	GGCGTTGATA	ATCTTTCTT	19920
TATCTTGAG	GAAGGTTTA	AAGACAGTCT	AAAAATAGG	ATGAACCTGC	TTAAGATTGT	19980
CCTCAATAAG	TCCGAAAAT	TTCTCTGGTT	CCTTATTCTG	GAAGTGAAAA	AGCAAGAGTT	20040
GATAGAGCTG	ATAGTGGTGT	TTCAAGTCTT	CCGAATAGCT	CAAAGCTTG	TTTAAAATCT	20100
CTTTATTGGT	TAAGTGCATA	CGAAAATAG	GACGATAAAA	TCGCTTATCA	CTCAGTTAC	20160
GGCTATCCTG	TTGAATGAGT	TTCCAGTAGC	GCTTGATAG			20199

(2) INFORMATION FOR SEQ ID NO: 7:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 19702 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:

ACCCGATGTA	TCAGCGGATA	TTTACTCTAT	TTTCACAAACG	ATGTTATAACC	CACAATAAAA	60
GAAAAAAAGAC	CCTAAGGTCT	CCTTTGCTTT	TATTATTTAAA	CGCGTTCAAC	TTTACCTGAT	120
TTCAAAGCAC	GAGCTGAAGC	CCAAACCTTT	TTAGGTTTAC	CATCGATAAG	AACAGTAAC	180
TTTTGAAGGT	TTGGTTTAC	GGCACGTTT	GTGTTGGTTCA	TCGCGTGTGA	ACGGTTGTT	240
CCTGATACAG	TCTTACGACC	TGTAAAGTAA	CATACTTTAG	CCATTGTGTT	TTCCCTCTAT	300
TAGATCTAAT	ATAGCGGATG	TGCTAGCACC	ACATACCGTA	CTATGTTATC	ACATTTCTT	360
GTGTTTTGCA	AGGAAATTGG	AAGATTTTTT	ATTTGTGTCT	TAAATCAGGT	CTTGCCTGAC	420
ATTTCTGCTC	TCCACATGCC	ATCGTTGATT	AACAGAACAC	CAGAATTAAA	ATTATGTGTA	480
TAAAAATCAT	CTCTAACTGC	AGCTAAGGGT	ATAGCCGTCA	AGTCCAAATC	CCACAGCTCA	540

195

TCTATCGATT TTCTTACAAC AATATCTGAA TCCAAATACA GTACACGAGA CTCGCTTACA	600
TACTTTGGAA TAAAATACCT AAAAAGCCG CATATGAAAG TCCCTCAAAG GGGAGACGAT	660
AACCTTCAG AATATTACTG TCAATCTAA CATTACAAT CTCACTATTCTAAAGTCTCTA	720
GTCTTTTTCATCAATTGG AACCAATTCTC GCGGAAGGTC ATCATTAAAA ACATAAAACT	780
TAAGATTATA ATGATGAACA CAAAGAGATT TTATTGTTGT TTCAACTTTA TCCATATAAG	840
CATTATCTGC ACCTAAGACA ATCGCTTTT TCTCTTCTT CACTTTTAT CTCATTTCTT	900
TTTATTCCCA TCATATTATT CCCATCATAT GTTTCCCATC ATATGTTCT ACGTAACCAT	960
TATTTTCGCC TATTGTTCG TAAAACCATA CCAGTGGAGA TTTTAGATGA AGTCCCATT	1020
CGGTTACAA TTTTACATT ACGACACGGA GTTTTACAAA TCGATTTCAT TTGCCAACG	1080
TAGTTAGTGA GGCAGTTAGC TAGTCGCCA AATAGCGACT AGCGTCCAAC AATTGGAAC	1140
TTTGTAGTCA ATTGTTGGTA CTGAGTCACA TCTTCTCCTC TAACTCTACG TCTGGATACT	1200
TGTCCGCAA CCACCGGAGG GCAAAGTCAT TTTCAAAGAG AAAGACTGGT TGGTCAAAC	1260
GGTCTTGGC TAAGATATTG CGACTTGACG ACATCCGTTCA ATCCAAGTCC TCAGGCTTGA	1320
TCCAACGAAC GGTCTTTTA CCCATTGGGT TCATAACTAC TTCCGCATTG TACTCGCCTT	1380
CCATGCGGTG TTTAAAGACT TCAAACGTGA GTTGACCTAC AGCGCCTAGC ATGTACTCAC	1440
CTGTTGGTA ATTCTTATAA AGCTGAACGG CTCCTTCTTG CACCAATTGC TCAATCCCCT	1500
TGTGGAGGA TTTTGCTTC ATAACATTCT TAGCAGAAAC TTTCATGAAA ATCTCAGGTG	1560
TAAAGGTTGG CAGGGTTCA AATTCAAAC TGTGTTTTCC AACCGTCAAG GTATCCCCAA	1620
CCTGATAAGT ACCGGTATCG TAAACCCCGA TAATATCACC TGCCACGGCA TTGGTCACAT	1680
TCTCACGACT CTCCGCCATA AACTGGTAA CATTAGATAG TTTAGCCCC TTACCAAGTAC	1740
GAGGGAGATT GACACTCATG CCGCGCTCAA ATTCGCCAGA TACGATACGG ACAAAGGCAA	1800
TACGGTCACG GTGACGAGGG TCCATGTTGG CTTGGATTTT AAAGACAAAG CCTGAGAAAT	1860
CCTTGTCTATA AGGATCCACA ATTCACCGT CTGTTTCTT GTGACCATGT GGTTCTGGAG	1920
CAAACTTGAG GAAGGTTCA AGGAAGGTCT GCACACCAAA GTTGTGAGG GCTGAACCGA	1980
AAAAGACAGG CGTCAATTCT CCAGCCAGAA TAGCTTCCCTC TGAAACTCA TTCCCGGCTT	2040
CATTTAAAG CTCAATGTCA TCCTTGACTT GCTCGTAGAA AGGATTGCTA CCAAAGAGTT	2100
TGTCCCCGTC TTCTAGACTG GCAAAACGCT CATCCCCTTT GTAAAGCTCT AACGTTGGT	2160
TATAGAGGTC ATACAAGCCC TCAAAGGCTT TCCCCATCCC GATAGGCCAG TTCATAGGGT	2220
AGCTAGCAAT GCCCAAGATT TCTTCCAATT CTTGCAAGAG ATCCAAAGGC TCACGACCGT	2280

196	
CACGGTCCAG CTTGTTCAT AAGGTAAGA CTGGAATGCC ACGATGTTTC ACAACCTCAA	2340
ACAATTCTT GGTTGAGCC TCGATCCCTT TGGCAGAGTC CACGACCAG ACCGCAGCAT	2400
CCACCGCCAT CAAGGTACGA TAGGTATCTT CTGAGAAGTC CTCGTCCCTT GGCGTGTCTA	2460
AGATATTACAC GCGCTTGCCG TCGTAGTCAA ATTGCATAAC AGATGAAGTA ACAGAAATCC	2520
CACGTTGCTT CTCGATATCC ATCCAGTCAG ATTTAGCAAA AGTCCCTGTT TTCTTCCCTT	2580
TTACCGTACCG AGCCTCACGA ATCTCACCCCC CAAAGTAGAG TAACTGCTCA GTGATGGTTG	2640
TTTTCCCCCGC GTCCGGGTGG GAGATAATGG CAAAGGTACG ACGTTTCTTA ATTTCTTCTT	2700
GAATATTCAAT AAGTTCTCTT TCTTTGATTC TCTATTTTC TTGTTCAAT AGCTGAGAAT	2760
GATTTTACA TTGGATTTA CCATTCCTTT CAACACTCCA TTATATCGGA TTTTAGCATT	2820
TTTTTCATT TCTATTTCTT TTCACCTCCC CCTCCCTTAT TTATAGGAAA ATATGGTAA	2880
ATAGAACAGA CTAAAATCA TCATTCACG AAAGGATGCA AGATGAAAAT TACGCAAGAA	2940
GAGGTAACAC ACGTTGCCAA TCTTCAAAA TTAAGATTCT CTGAGAAGA AACTGCTGCC	3000
TTTGCACCA CCTTGTCTAA GATTGTTGAC ATGGTTGAAT TGCTGGCGA AGTTGACACA	3060
ACTGGTGTG CACCTACTAC GACTATGGCT GACCGCAAGA CTGTAACCCG CCCTGATGTG	3120
GCCGAAAGAAG GAATAGACCG TGATCGCTTG TTTAAAAACG TACCTGAAAA AGACAACCTAC	3180
TATATCAAGG TGCCAGCTAT CCTAGACAAT GGAGGAGATG CCTAATGACT TTTAACAAATA	3240
AAACTATTGA AGAGTTGCAC AATCTCCTTG TCTCTAAGGA AATTCTGCA ACAGAATTGA	3300
CCCAAGCAAC ACTTGAATAT ATCAAGTCTC GTGAGGAAGC CCTCAATTCA TTTGTCACCA	3360
TCGCTGAGGA GCAAGCTCTT GTCAAGCTA AAGCCATTGA TGAAGCTGGA ATTGATGCTG	3420
ACAATGTCCTT TTCAGGAATT CCACCTGCTG TTAAGGATAA CATCTCTACA GACGGTATTTC	3480
TCACAACCTGC TGCCCTAAAA ATGCTCTACA ACTATGAGCC AATCTTGAT GCGACAGCTG	3540
TTGCCAATGC AAAAACCAAG GGCGATGATTG TCCTGGAAA GACCAACATG GACGAATTG	3600
CTATGGGTGG TTCAGGTGAA ACTTCACACT ACGGAGCAAC TAAAAACGCT TGGAAACCACA	3660
GCAAGGTTCC TGGTGGGTCA TCAAGTGGTT CTGCCGCAGC TGTAGCCTCA GGACAAGTTC	3720
GCTTGTCACT TGGTTCTGAT ACTGGTGGTT CCATCCGCCA ACCTGCTGCC TTCAACGGAA	3780
TCGTTGGTCT CAAACCAACC TACGGAACAG TTTCACGTTT CGGTCTCATT GCCTTGGTA	3840
GCTCATTAGA CCAGATTGGA CCTTTGCTC CTACTGTTAA GGAAAATGCC CTCTTGCTCA	3900
ACGCTATTGC CAGCGAACAG GCTAAAGACT CTACTTCTGC TCCTGTCCGC ATCGCCGACT	3960
TTACTTCAAA AATCGGCCAA GACATCAAGG GTATGAAAAT CGCTTGCCT AAGGAATACC	4020
TAGGCGAACAGG AATTGATCCA GAGGTTAAGG AAACAATCTT AAACGCGGCC AAACACTTTG	4080

AAAAATTGGG TGCTATCGTC	GAAGAAGTCA	GCCTTCCTCA	CTCTAAATAC	GGTGTGCCG	4140
TTTATTACAT CATCGTTCA	TCAGAAGCTT	CATCAAACCTT	GCAACGCTTC	GACGGTATCC	4200
GTTACGGCTA	TCGCGCAGAA	GATGCAACCA	ACCTTGATGA	AATCTATGTA	4260
GCCAAGGTTT	TGGTGAAGAG	GTAAAACGTC	GTATCATGCT	GGGTACTTTTC	4320
CAGGTTACTA	TGATGCCTAC	TACAAAAAGG	CTGGTCAAGT	CCGTACCCCTC	4380
ATTTCGAAAA	AGTCTTCGCG	GATTACGATT	TGATTTGGG	TCCAAGTGCCT	4440
CCTATGACTT	GGATTCTCTC	AACCATGACC	CAGTTGCCAT	GTACTTAGCC	4500
CCATACTGT	AAACTGGCA	GGACTGCCCTG	GAATTTGAT	TCCTGCTGGA	4560
GTCTACCTGT	CGGACTCCAA	TTGATTGGTC	CCAAGTACTC	TGAGGAAACC	4620
CTGCTGCTGC	TTTGAGCA	ACAACAGACT	ACCACAAACA	ACAACCCGTG	4680
GTGACAACTA	ATGAACATTG	AAACAGTCAT	CGGACTGAA	GTCCACGTTAG	4740
CAATTCAAA	ATCTTCTCAC	CTACTTCTGC	CCACTTTGGA	AATGACCAAA	4800
TAACGTGATT	GACTGGTCTT	TCCCAGGAGT	TCTACCAGTT	CTCAATAAAG	4860
TGCCGGTATC	AAGGCTGCTC	TTGCCCTCAA	CATGGACATC	CACAAAAAGA	4920
CCGCAAGAAC	TACTTCTATC	CTGATAACCC	CAAAGCCTAC	CAAATTCTC	4980
ACCAATCGGA	TATAATGGCT	GGATTTGAAGT	CAAACATAGAA	GACGGTACGA	5040
CGGTATCGAA	CGTCCCCACC	TAGAGGAAGA	CGCTGGTAAA	AAACACCCATG	5100
CTACTCTTAT	GTGACCTCA	ACCGCCAAGG	GGTCCCTTGT	ATTGAGATTG	5160
AGATATGCGT	TCTCCTGAAG	AAGCCTATGC	TTATCTGACA	GCCCTCAAGG	5220
GTACGCTGGC	ATTTCTGACG	TTAAGATGGA	GGAAGGTTCG	ATGCGTGTGG	5280
CTCCCTTCGT	CCTTATGGTC	AAAGAGAAATT	CGGTACCAAG	ACTGAATTGA	5340
CTCCTCTCA	AACGTTCGTA	AAGGTCTTGA	ATACGAAGTC	CAACGCCAGG	5400
TCGCTCAGGT	GGTCAAATCC	GCCAAGAAC	ACGCCGTTAC	GATGAAGCGA	5460
CATCCTCATG	CGTGTCAAGG	AAGGGGCTGC	TGACTTACCGC	TACTTCCAG	5520
ACCCCTCTTT	GAATTTCTG	ACGAGTGGAT	TGAGGAAATG	CGGACTGAGT	5580
TCCAAAAGAA	CGTCGTGCGC	GTATGTATC	TGACCTTGTT	TTATCAGACT	5640
TCAGTTGACT	GCTAATAAAG	TCACTTCTGA	CITCTTTGAA	AAAGCTGTTG	5700
TGATGCCAAA	CAAGTCTCTA	ACTGGCTCCA	AGGGGAAGTC	GCTCAGTTCT	5760
AGGTAAAACA	CTGGAACAAA	TCGAATTGAC	ACCGAAAAAC	TTGGTTGAAA	5820

198	
CATCGAAGAC GGTACTATTT CATCTAAGAT TGCCAAGAAA GTCTTGTCC ATCTAGCTAA	5880
AAATGGCGGT GGCGCGCGTG AATACGTGGA AAAAGCAGGT ATGGTCAAA TTTCAGATCC	5940
AGCTATCTTG ATCCAATCA TCCACCAAGT CTTTGCCTG AACGAAGCTG CTGTTGCCGA	6000
CTTCAAGTCA GGCAACAGTA ACAGCGACAA GGCTTTACAG GATTCCCTAT GAAGGCAACC	6060
AAAGGCCAAG CCAACCCACA AGTTGCCCTT AAAACTACTTG CACAGGAATT GGCGAAGTTG	6120
AAAGAAAAGT AGACAGAACAA AAACCAAGCCC TAAGGTTGGT TTTTCTTCT CTACCAACTC	6180
CCAATAACTA TTTTGGCTTT ATTTCCAGAG TATTTTATGG TAAAATGAAG AGTAATAATA	6240
TTTATTAAG AGGTAAAAAC ATGATTGAAG CAAGTACCTT AAAAGCTGGT ATGACCTTG	6300
AAACAGCTGA CGGCAAATG ATTGCGTTT TGGAAGCTAG TCACCACAAA CCAGGTAAG	6360
GAAACACGAT CATGCGTATG AAATTGCGTG ATGTCCTACG TGGTTCTACA TTTGACACAA	6420
GCTACCGTCC AGAGGAAAAA TTTGAACAAG CTATTATCGA GACTGTCCCC GCTCAATACT	6480
TGTACAAAAT GGATGACACA GCATACTTCA TGAATACAGA AACTTATGAC CAATACGAAA	6540
TCCCTGTAGT CAATGTTGAA AACGAATTGTC TTTACATCCT TGAAAACCTCT GATGTGAAA	6600
TCCAATTCTA CGGAACGTGAA GTGATCGGTG TCACCGTTCC TACTACTGTT GAGTTGACAG	6660
TTGCTGAAAC TCAACCATCT ATCAAAGGTG CTACTGTTAC AGGTTCTGGT AAACCAGCAA	6720
CGATGGAAAC TGGACTTGTC GTAAACGTTC CAGACTTCAT CGAACGAGGA CAAAAACTCG	6780
TTATCAACAC TGCAGAAGGA ACTTACGTTC CTCGTGCCTA ATCTCTAGAA AGAGGTCATT	6840
CTATGGAAT TGAAGAACAA CTTGGCGAAA TCGTTATCGC CCCACGTGTA CTTGAAAAAA	6900
TCATTTGCTAT CGCTACTGCA AAGGTAGAGG GTGTTCACTC TTTTTCAAAC AGATCAGTGT	6960
CTGATACCCCT TTCAAAACTT TCACTCGGCC GTGGCATTTA TCTTAAAAAC GTGGACGAA	7020
AACTCACAGC AGATATCTAT CTCTACCTTG AGTACGGAGT AAAAGTTCCCT AAGGTAGCGG	7080
TTGCTATCCA GAAAGCTGTC AAAGATGCCG TCCGTAATAT GGCTGATGTA GAACTCGCTG	7140
CTATCAATAT TCACGTTGCA GGTATCGTCC CAGATAAAAC ACCAAAACCA GAATTGAAAG	7200
ATCTATTGTA CGAGGACTTC CTCAATGACT AGTCCACTAT TAGAATCTAG ACGCCAACCTC	7260
CGTAAATGCG CTTTCAAGC TCTCATGAGC CTTGAGTTCG GTACGGATGT CGAAACTGCT	7320
TGTCGTTTCG CCTATACTCA TGATCGTGA GATACGGATG TACAACCTCC AGCCTTTTG	7380
ATAGACCTCG TTTCTGGTGT TCAAGCTAAA AAGGAAGAAC TAGATAAGCA AATCACTCAG	7440
CATTTAAAAG CAGGGTGGAC CATTGAACGC TTAACGCTCG TGGAGAGAAA CCTCCTTCGC	7500
TTGGGAGTCT TTGAAATCAC TTCATTGAC ACTCCTCAGC TGGTTGCTGT TAATGAAGCT	7560
ATCGAGCTTG CAAAGGACTT CTCCGATCAA AAATCTGCCG GTTTTATCAA TGGACTGCTC	7620

199

AGCCAGTTG TAACAGAAGA ACAATAAGGC TCTTTGTCAA CTGTAGTGGG TTGAAAAAAA	7680
GCTAAGCTCG AGAAAGGACA AATTCGTCC TTTCTTTTT GATGTTCAA GCGATAAAAA	7740
TCCGTTTTT GAAGTTTCGA AAGTTTCGAA ACCAAAGGC ATTGCGCTTG ATAAGTTGCA	7800
TGAGATTATT GGTGCGCTCC AGTTGGCAT TAGAATAGTG TAGTTGAAGG GCGTTGACAA	7860
TCTTTCTTT ATCTTGAGG AAGGTTTAA AGACAGTCTG AAAATAGGA TGAGCCTGCT	7920
TAAGATTGTC CTCAATAAGT CCGAAAAATT TCTCTGGTC CTTATTCTGG AAGTGAAACA	7980
GCAAGAGCTG ATAGAGCTGA TAGTGGTGT TCAAGTCTTG TGAATGGCTC AAAAGCTTGT	8040
CTAAAATCTC TTTATTGGTT AAGTGCATAC GAAAAGTAGG ACCATAAAAT CGCTTATCAC	8100
TCAGTCTACG GCTATCCTGT TGAATGAGTT TCCAGTAGCG CTTGATATCC TTGTATTCTAT	8160
GGGATTTCG ATGAAACTGA TTCATGATTT GGACACGCAC ACGACTCATG GCACGGCTAA	8220
GATGTTGTAC AATGTGAAAG CGATCAAGAA CGATTTAGC ATTGGGGAGT GAAACAGTCT	8280
GGGAGACTGT TTCAGCCTGA GCCTAGGAAT TTGAAAGCGA AGCTGTTAG CCAAGTCATA	8340
GTAAGGGCTA AACATATCCA TAGTAATAAT TTGACCGCA CATCGGACAA CTCTATCGTA	8400
GCGAAGAAAG TGATTCGAA TGATAGCTTG TGTCTACCC TCAAGAACAG TGATGATATT	8460
GAGATTGTTA AAATCTTGCG CAATGAAGCT CATCTTCCC TTTGAAAAG CATACTCATC	8520
CCAAGACATA ATCTCAGGAA GACAAGAAAA ATCATGTTA AAGTGAAAT CATTGAGCTT	8580
ACGAATAACA GTTGAAGTTG AGATGGAAAG CTGATGGCA ATATCAGTC TAGAAATCTT	8640
TTCAATCAAC TTTGAGCAA TCTTTGGTT GATGATACGA GGGATTTGGT GATTTTCTT	8700
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TCTAAGGAGA ATTCTAGTAG CCATACCAGT CGTTCAAGA TAAGGAATT TAGAAGGTTT	8820
TTGAAAGTCA TATTCTTCA ATTGGTTTCC GCACTCAGGG CAAGATGGGG CGTCGTAGTC	8880
CAGTTGGCG ATGATTCCT TGTGTGTATC CTTATTGATG ATGTCTAAA TCTGGATATT	8940
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AAAGAAAAAG TGTTGATAG ATATCAAACA CTTTTTCTT TGCCTCCAC TATCTAAAAA	9180
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CAAAAGAA ACCAATCAGA CTATAATATA ATAAACTAAT TGGATCTCTG TGAGATACTA	9360

200	
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TGATTACAGG CAGGACAAAA GATAAGATAG TCGATAAAAA GGTTGGTTGT CCATTTGAAA	9540
AAAGCACGGT AAAATACTCA TCATGAATAT TCCTATGATT AATCAAATGA GCATAGCGTG	9600
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ACCAGTTCCA GCTCTTTTC TCAAAGATAA AGAGCATCTT TTTCTTTTT AACCTCCAAA	9720
TTAATAGAAG GAAACCTCCC ACTAATCCCA TTGTTAAAAT AAGAGAATAG ACATCAGCTC	9780
CTAACCTAA AATGATCGTC ACATACAATC CAATTGTTG TGGTAAATAG GTAGATAGTA	9840
AAATAATAAG CAAAAATATT CCAAATTGTC TTAGTTTTT TGTGTTCTC ATCGTACTTT	9900
TTTGAAAGAT TACCTGCTC GGAAGCCGTA CTTCCAAGCA TCTATATAAG AATTAAGTGC	9960
CCCTTGCTC ATATAGGGAG CAAATTCTCT ATAATATAAC CATCTACTAT ATCCATCTTC	10020
CCAAACAGCA AGACCACCTG AAGTTTGCTC CAAGTCCTCA GTTGAAGAA CTGTAATGT	10080
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GTATATTCTTCTT TTTTTGCTC ATTTTATAGC CCATCTCCTC AACTGGCAAT TTTTCGACCT	10200
GAATTACATT TTTCCATAAA AAATGAGACC TTTCTAGTCT CATTAGTCA TTCTTAGTAT	10260
TTTCTAAATC GTTGATAGCG TTCTTCCAGC AACTCTTCTA GCGGTTTTG TGAAAGTCTA	10320
GCCAGCTCG TTTGGAGTTC TTTTTGACA CTCTTAATCA GTTCTTTACT AGAAAGTCCT	10380
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ATCAGTTCTG CTGCTTCCAT AGCGCGAGTA CCGTCCTTCC ATAAAATGGA AGCAAAGCCT	10500
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AGAGCCAGAG CCCCGCCTGA ACCACCTTCA CCGATAATAA TGGCGATAAT AGGAACTTTC	10620
AGGTCACTCA TTTCATGAG ATTGCGAGCG ATAGCTTCCC CTTGACCACG TTCTTCCGCT	10680
CCGACACCAAG GATAAGCACC TGCTGTATTG ATAAAGGTCA CAACTGGACG GCCAAATTTC	10740
TCAGCCCTGTT TCATCAACCG CAGTGCCTTT CGGTAGCCTT CTGGATGTGG TTGGCCAAA	10800
TTCCGTTGA GGTTGTCTTG CAAACTCTTG CCTTTTGGA TACCAACCAC TGTTACAGCT	10860
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CCATGTAATT GGATAAAATTG ATCAAAATG CCTGTCGCAA AGTCCAAGGT TGTCAAGCGA	10980
CTCTGCTCAC GCGCTCTCT GACTATTTT GCAATATTCA TCTAGGACTC CCTCCATGCA	11040
ATCTGACTAG GCTAGCAATC GTATCTGGTA AGTCTCTTCTT TTTGACAATA GCATCCACAA	11100
AGCCATGTTCA TAATAGGAAT TCTGCCTTTT GGAAATCCTC AGGCAAGCTT TCACGAACCG	11160

201

TATTTTCAAT	CACACGACGC	CCAGCAAAAC	CAACCAAGCT	CTGTGGTTCA	GCCAGAATGA	11220
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TCAGGTAAAA	GAGACCAGCA	TTTGAATGGC	GTAAACCGC	CGCAGAGATC	TTAGCCATCT	11340
GCATGAGACT	CATGATTCT	TCCTGCATAC	GGGCTCCACC	AGAGGCTGTG	AATAGGACAA	11400
CTGGCAATTT	TTCGACAGTC	GCATACTCAA	ACAAACGAGT	GATTTTTCA	CCTACAACCG	11460
TACCCATAGA	AGCCATGATA	AAGTTAGAAT	CCATAATCCC	AAGAGCCACA	GTCTGACCTT	11520
TAATAAGAGC	AGTTCCGTGTC	ACAACGGCTT	CATGCAGACC	TGTTTTTCA	CCCATAGATG	11580
CCAGTTCTT	TTGGTAACCA	GGGAATGCA	AGGGATCCTT	GCTTCATC	CCTGTAAACA	11640
ATTCTTGAA	GGTTCCCATCA	TCAATCGTCA	AAGCCAAGCG	TTCTTGGCA	GAAATACGAA	11700
AGGTATAGCT	ACAGTGCAGA	CAGATACTT	CACTTCCAG	ATCCTTCTGA	TAGATGGTAT	11760
GCTTACAGCC	TGGACACTGG	GAAAATAATT	CATCTGGAAC	CTCTGGCTTA	GCTTGAGGTT	11820
TTTCCCTAAC	CGAACGATTG	GGATTGATTC	GAATATACTT	ATCTTTTTA	CTAAATAGAG	11880
CCATTGATTC	CCCTTTCCG	TTTAAACTCT	TAAAGTCATT	TTATTCTTT	TCTTGATATT	11940
TAGGTAAGAA	GGTTCCATC	AAGAAGGAAG	TATCATAATC	CCCAGCAATG	ACATTGCGAT	12000
CTGAAATGAG	GTCAAGCTGG	AAATCTGCAT	TGGCTGAC	TCCTCAATT	TCTAATTCAT	12060
AGAGGGACG	TTGCATTTTC	ATCAAGGCAGT	CAAACGATT	TCGCGCGTGT	ACTATGATT	12120
TGGCAATCAT	ACTATCATAA	TAAGGCGGAA	TGGTATAACC	TGGTAAACT	GCTGAATCCA	12180
CGCGCAAGCC	AACTCCACCA	CTTGGCAGAT	AGAGATTAGT	AATCTTACCT	GGACTTGGAG	12240
CAAAGTTAAA	GGCTGGTTT	TCTGCATTGA	TACGACACTC	GATGGCATGA	CCCGCGTAGGA	12300
CAATATCTTC	TTGCTTAACA	GACAAAGGCT	GACCTGCCGC	AATGCAATC	TGTTCCCTAA	12360
CGATATCAAC	ACCTGAAACA	AACTCTGTTA	CTGGATGTT	TACCTGAACA	CGAGTATTCA	12420
TCTCCATGAA	ATAGAAATTG	CTACTTGCTT	CATCAAGAAG	AAATTCAATG	GTTCCCTGCAT	12480
TCTCATAGCC	AACAAACTCT	GCCGCTCGAA	CAGCAGCAGC	ACCTATTCA	TGACGCAGCG	12540
TTTTCCGAT	TGCAATCGAG	GGACTTTCTT	CCAAAACCTT	TTGGTTATTC	CTTGAAAGAG	12600
AACAATCCCG	TTCACCCAAG	TGAATCACAT	GTCCATGCTC	ATCACCTAGG	ATTTGAACCT	12660
CAATGTGCCG	AGCTGGATAG	ATAACCCGTT	CTATGTACAT	GGCACCATTG	CCATAATTGG	12720
CCTTGGCCTC	ACTAGAGGCA	GTTCAAAAGG	CAGAAACGAG	GTCATCTGGT	TTTTCAACCT	12780
TACGAATCCC	TTTACCCACCT	CCACCTGCTG	AAGCCTTGAG	CATAACAGGA	TAGCCAATT	12840
TTTCAGCAAC	AATCAAAGCT	TCTTCAGAGT	TATGCACTTC	TCCATCTGAA	CCTGGTATAA	12900

202

CAGGCACACC	TGCTTTAAC	ATCTGAGCAC	GCGCATTGAT	CTTATCCCC	ATCATATCCA	12960
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TGGAATTTC	ACTGAGAAAT	CCAAAACCG	GGTGAATAGC	TTCTGCCTCA	GTCAAGACTG	13080
CAGCTGATAG	AACTGCATTA	ATATTGAGAT	AAGACTCTGT	TGCCTTGCCA	GGACCAATAC	13140
AAACTGCTTC	ATCTGCCAAA	AGCGTATGAA	GAGCTTCCTT	ATCAGCAGTT	GAATAAACCG	13200
CTACCGTCGC	AATCCCCAAT	TCACGTGCCG	CACGGATAAT	ACGAACCGCA	ATTTCACCCAC	13260
GATTGGCAAT	TAAAATTTT	CGAAACATGG	AGAACCTCCT	TAGTTCCCAA	TTGCAAAAGT	13320
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GGTGCCACGA	CGTTTACAA	AAGTCGCTGT	CATAACCAAT	TGGTCGCCTG	GTACAACATTG	13440
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CATAACTGGG	TATTGAGGAA	AGTGGCCGTT	AAAGAAAGGC	TCGTTGATGG	TCACATTTT	13620
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TTACCAAAC	CAACCATTTC	TTCGTTAGAG	ACGAGAATT	CCGTTACCAC	ACCATCCTTA	13800
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ACTCCAACAA	GTGGACTCTC	TACAAGATTT	CCCTCAGTAG	CCACACTTGC	TTCAGCTGGA	13980
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GTTGCTAGAA	CGGGTGCTGG	AGGCAGTGA	GTTGCAACTT	CAGGCACAGG	TCTTGCTTCA	14100
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CTTCACATTGT	CCCAGCTGTC	ATTGGTACAA	AGTTATGACG	CATAGCTTCG	ATGGTGACGA	14400
TAGCTTCTAC	TGCACCCGCA	GCCCCCAGCA	AATGTCCTGT	AAAAGACTTG	GTTGATGATA	14460
CAGGTACTTC	CTTACCAAGA	ACAGCTACGA	TAGCACCAC	TTCTCCTTTT	TCATTGGCAG	14520
GAGTTGACGT	TCCGTGAGCA	TTGACATAGG	CTACTTGCTC	TGGAGAAATC	TCAGCTTCTT	14580
CCAAGGCTAG	TTTGATGGCC	TTGATAGCTC	CCTGACCTTC	TGGATGTGGA	GAAGTCATGT	14640
GGTAGGCATC	ACAAGTATTT	CCGTAACCAA	CCACTTCAGC	CAGGATAGTA	GCTCCACGTT	14700

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CATTGCGATC	CTTATCAAAT	GGGATCGAAG	CACGAGTTGG	ATCCTCTGTA	GTAGAGAGAG	14820
CTGTTAACGC	TTGGAAACCA	GGCATGGCAA	AAGGTGTGAT	AGAAGCTTCT	GTTCCCTCCCA	14880
CCAACATCAC	ATCTTGAAA	CCAAACTTAA	TGGAGCGGAA	GGCATCCCCA	ATCGCATCAT	14940
TTGATGAAGA	GCAGGGACTA	TTGATAGATT	TACAAACACC	GTTTGCACCA	AAACGCATGG	15000
CTACATTCCC	AGAAGCCATA	TTTGGTAAAG	CTTTTGGAG	AGTCATTGGT	TTGACACGTT	15060
TGGGTCCTTT	TTCATGAAGG	CGAAGTACCT	GATCTTCAAT	TTCCTTGATT	CCACCAATAC	15120
CAGATGCAAC	GATAACACCA	AAACGATCCC	TATTAAGAGC	CTCTACATCA	AGATTGGCAT	15180
GATTTACAGC	CTCTTGGCT	GCATACAAGG	CATATAAAAGA	ATAGTTATCA	AAACGGTTGG	15240
TATCTTTTT	TACAAAGTAT	TTATCGAACG	GAAAATCTTG	GATTTCTGCC	GCATTATGCA	15300
CATCAAAGTC	ACTATGATCA	AATTTTGTA	TGCCACCAAT	GCCGATTTTC	CCAGTTGCTA	15360
AACTATTCCA	AAATTCTCT	GGTGTATTTC	CGATTGGAGA	TGTTACTCCA	TAACCTGTTA	15420
CCACTACTCG	ATTTAGTTTC	ATTCTTTCA	CCTCTAGCTT	TCGCTACATA	CTTAAGCCAC	15480
CATCAATGGC	AACCACCTGT	CCAGTTAGAT	AATCTTGGCC	TGCTAAAAAT	ACTGTCAAAT	15540
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TGACTCGTAT	ATTCCGACTA	GCGACCTCGC	GTGCCACAGA	CTTGGTAAAG	CCAAATCACCC	15720
CAGCCTTAGA	AGCAGCATAA	TTAGCTTGAC	CAATATTCCC	CATCAAACCA	ACAAACACTAG	15780
ACATATTAAT	GATAGCACCT	TCTCTGGCTT	TCATCATCGG	TTTCAAGACT	GATTGTGTCA	15840
TATTAAGGC	ACCAGTCAGA	TTGACCTTGA	GCACCTTTTC	AAAATCTGCT	TCTGTCATCT	15900
TGAGCATAAG	AGTATCTTGG	GTAATCCCTG	CATTGTTGAC	AAAACACATCT	ACTGAACCCA	15960
GTTCTGCAAT	AGCTTGATCA	ATCATAACGCT	TAGCGTCTGC	AAAATCTGAT	ACATCTCCTG	16020
AAATGGGAAC	CACCTTGATA	CCATAGTTTG	AAAACTCAGC	GAGCAATTCT	TCTGAGATTG	16080
CCCCACGACT	GTAAAGACA	ATGTTGGCTC	CTGCTTGAGC	AAACTTGTGG	GCGATGGCAA	16140
GACCAATTCC	ACGACTCGAA	CCTGTAATAA	AGATATTTTT	ATGTTCTAGT	TTCAATTCTT	16200
TCCTTCTAAA	ACTTCTACTT	ATTTTAGTCT	ATTTTCTAA	AAAGTGTACT	AAACTCGCTT	16260
GATCTTCCAC	ATGAGCTAAG	TGAGCAGTTT	GATCAATTTC	TTAACAAAA	CCTGACAAGA	16320
CTTTCCCCGG	TCCAATCTCG	ATAAAAGTTGC	TTATGCCTGC	TTCTTGCATG	ACCCCAATAC	16380
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204

TTTGCATCAC	AGCAGCTTCT	GTATTGCCGA	CTAGGGGACA	AGTAAAATCT	GAAAAACTTA	16500
CCTGAGCTAG	AGTTTCAGCT	AGTTTCTGGC	TAGCAGGTTC	AAGGAGAGCG	GTGTGAAAGG	16560
GACCTGACAC	CTTAAGAGGA	ATCAAGCGTT	TGGCACCTGC	TTCTTGCAAA	AGTTCAACCG	16620
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TCAAAGATTT	CTAAATCAGG	ATCTTCTTGC	TTAAAGGCAT	CTTTTCAGC	CAGTTCAAAA	17400
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TCTTTGCAA	CTACAAACCG	TGTCCCCACC	TGTACAGCCT	CTGCACCTAG	CATAAAGCCA	17580
GCCGCAGCAC	CTTCACCAC	CGCAATTCC	CCTGCAGCAA	TAACAGGAAT	AGATATAGCT	17640
GTGGCTACCT	GTGCCACCAA	GGTCATGGTT	GTAAATTAC	CGATATGCC	CCCAGCTTCC	17700
ATTCCTCTG	CAATAACAGC	GTCTGCACCG	ATTTTTCCA	TGCGTTAGC	AAAGGCAC	17760
CTAGGAACAA	CAGGAATAAC	GATTATCCC	GCTTCATGGA	AACGTTCCAT	ATACTTGCTT	17820
GGATTTCTG	CTCCTGTTGT	GACAACTTTA	ACACCTTCTT	CAATAACGAG	ATCCACGATG	17880
TCTTCCACAA	AGGGAGATAA	GAGCATGATG	TTGACCCCAA	AGGGTTTATC	AGTCAATGAT	17940
TTGATTTTAT	CAATATTGGC	CTTGACAAC	TCTTCCGGGG	CATTTCCCC	ACCGATAATT	18000
CCTAATCCCT	CAGCCTTGGA	AACAGCCCC	GCCAAATCAC	CATCAGCAAC	CCAGGCCATC	18060
CCTCCTTGGA	AAATAGGATA	ATCAATCTTC	ATAAATTCTG	TAATACGCGT	TTTCATAGTG	18120
CCTCCAACCT	TCCTTGCTTA	CGTAATAGTT	CGATTTCAC	ATAATTGAC	AGTCACAACTA	18180
TTACCTAAAC	AAGAGGGAGT	GGGTTCTCC	CTACTCCTTC	TACTAATATT	CTGCTTATT	18240

205

TGCTTGCTCT	TCAACGTAAG	CAACCAAGTC	ACCAACTGTT	TTCAAGTCAT	TTTCTGCTTC	18300
GATTTGGATA	TCAAAAGCAT	CTTCGATTTC	TGAGATTACT	TGGAACAACT	CCAATGAATC	18360
TGCGTCCAAA	TCATCAAAAG	TTGATTCAAG	TGTTACTTCT	GATGCGTCTT	TTCCAAGTTC	18420
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ATAGTTTTTT	TATAACAATG	TGTTCACCC	ATGATTACCT	AAATTGTAAG	AATGAGCGTG	18540
CCCCAGGTCA	AGCCTCCACC	GAAGCCTGAT	AGAAGAACAG	TCTGGCTACC	ATCTAAAGGG	18600
ATGAGACCTT	GTTCTACACA	CTCTGAAAGT	AAAATCGGGA	TACTGGCTGC	ACTGGTATTG	18660
CCATATTCCA	TCATAATTGCC	TGGAAGTTTG	GCTCGGTCAA	CACCAATTTC	TCTAGCCATC	18720
TTATCCAAA	TACGGTCATT	GGCTTGATGA	AGTAGCAGAT	AATCCAAGTC	TGTCACCTCT	18780
ATAGGAGATT	CATCAATAGT	CTGCTTGATA	GACTTGGCTA	CATCTCGAAT	GGCAAAATCA	18840
AAGACTGTGC	GTCCATCCAT	CTTCAAAAC	GAATCTGCAC	TTTCTTGATC	TGAAAATGGA	18900
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CCAAACAAACA	CAGCTGTTGA	TCGATCCGAC	CAATCGACTG	CCTTAGAGAG	GGTTTCAC	19080
CCAATCACCA	AGCCCTTTTG	AAAGCGACCA	GAAGCGATAA	ACTTTTCAGC	AGTTGAAAGA	19140
GCAAATACAA	ATCCACTGCA	AGCCGCGGTT	AACTCAAAG	CAAAGGCTTT	ATTAGCACCA	19200
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GTTCGACTTG	AAATCCACTC	ATCATTGGTA	TCCATAATCT	GAGCCAAGTC	GTGATTGTA	19440
ACCACTTGCT	CTGGCACATA	ATGAGCAACC	TGACTTATTT	TTGCAAAAGC	CATTATTC	19500
AATCCCTCAA	AAATTGGTAA	AGATTAGTCA	AACCTTAC	CATGACAGCA	ATTCTTC	19560
CGCTCATGCC	ATCAATAATT	TTTCTACCA	TGGCCTTG	GAAGCGTTA	TGCAGTCTAT	19620
GAATCAAGCG	ACCCCTCTT	GTCAAATGCA	GATGCCAAC	ACGACGATCC	TGTTCTGACC	19680
GAACCTCGCTC	AATGTAGCCCC	GG				19702

(2) INFORMATION FOR SEQ ID NO: 8:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6211 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:

GAAAATTTC	TCTCTTCTCT	TGAAAAAATTT	TGAAAAAAATG	GTATGATAGT	AACAAGTTAT	60
TTTTAAGAGG	AAAGAAAGGG	GAATAATGGA	GAAAATCAGT	TTAGAATCTC	CTAAGACGGG	120
GTCGGACCTA	GTTTGGAAA	CACTTCGTGA	TTTAGGAGTT	GATACCATCT	TTGGTTATCC	180
TGGTGCG	GTGTTGCTT	TTTATGATGC	GATATATAAT	TTTAAAGGC	TTGCCACAT	240
TCTAGGGCGC	CATGAGCAAG	GTTGTTGCA	TGAAGCTGAA	GGTTATGCCA	AATCAACTGG	300
AAAGTTGGGT	GTTGCCGTG	TCACTAGTGG	ACCAAGGAGCA	ACAAATGCCA	TTACAGGGAT	360
TGCGGATGCC	ATGAGCGATA	GCGTTCCCCT	TTTGGTCTTT	ACAGGTCAGG	TGGCGCGAGC	420
AGGGATTGGG	AAGGATGCC	TTCAGGAGGC	AGACATCGTG	GGATTACCA	TGCCAATCAC	480
TAAGTACAAT	TACCAAGTTC	GTGAGACAGC	TGATATTCCG	CGTATCATTA	CGGAAGCTGT	540
CCATATCGCA	ACTACAGGCC	GTCCAGGGCC	AGTTGTAATT	GACCTACAA	AAGACATATC	600
TGCTTTAGAA	ACAGACTTCA	TTTATTCAACC	AGAAGTGAAT	TTACCAAGTT	ATCAGCCGAC	660
TCTTGAGCCG	AATGATATGC	AAATCAAGAA	AATCTGAAG	CAATTGCTCA	AGGCTAAAAA	720
GCCAGTCITG	TTAGCTGGTG	GTGGAATTAG	TTATGCTGAG	GCTGCTACGG	AACTAAATGA	780
ATTTGCAGAA	CGCTATCAAA	TTCCAGTGGT	AACCAGTCTT	TTGGGACAAG	GAACGATTGC	840
AACGAGTCAC	CCACTCTTTC	TTGGAATGGG	AGGCATGCAC	GGGTCAATTG	CAGCAAATAT	900
TGCTATGACG	GAAGCGGACT	TTATGATTAG	TATTGGTTCT	CGTTTCGATG	ACCGTTTGAC	960
GGGGAACTCCT	AAGACTTTCG	CTAAGAATGC	TAAGGTTGCC	CACATTGATA	TTGACCCAGC	1020
TGAGATTGGC	AAGATTATCA	GTGAGACAT	TCCTGTAGTT	GGAGATGCTA	AGAAGGCCTT	1080
GCAAATGTTG	CTAGCAGAAC	CAACAGTTCA	CAACAACACT	AAAAAGTGG	TTGAGAAAGT	1140
CACTAAAGAC	AAGAATCGTG	TCGTTCTTA	TGATAAGAAA	GAGCGTGTGG	TTCAACCGCA	1200
AGCAGTTATT	GAACCAATTG	GTGAATTGAC	GAATGGAGAT	GCCATTGTGG	TAACAGACGT	1260
TGGTCAACAC	CAAATGTGGA	CAGCTCAGTA	TTATCCTAC	CAAATGAAC	GTCAGTTAGT	1320
GACTTCAGGT	GGTTGGGAA	CAATGGCCTT	TGGAATTCCA	GCAGCAATCG	GTGCTAAAAT	1380
TGCTAACCCA	GATAAGGAAG	TAGTCTGTT	TGTTGGGGAT	GGTGGTTTCC	AAATGACCAA	1440
CCAGGAGTTG	GCTATTTGA	ATATTTACAA	GGTCCAATC	AAGGTGGTTA	TGCTGAACAA	1500
TCATTCACTT	GGAATGGTTC	GCCAGTGGCA	GGAATCCTTC	TATGAAGGCA	GAACATCAGA	1560
GTCGGTCTTT	GATAACCTTC	CTGATTCCA	ATTGATGGCG	CAGGCTTATG	GTATTAAGAA	1620
CTATAAGTTT	GACAATCCTG	AGACCTTGGC	TCAAGACCTT	GAAGTCATCA	CTGAGGATGT	1680

TCCTATGCTA ATTGAGGTAG ATATTTCTCG TAAGGAACAG GTGTTACCAA TGGTACCGGC	1740
TGGTAAGAGT AATCATGAGA TGTGGGGGT GCAGTTCCAT GCCTAGAATG TAAACAGCAA	1800
AACTACAAAA TCCTTCAGGA GTCCTCAATC GCTTACAGG TGTCTATCT CGTCGTCAGG	1860
TTAATATTGA AAGCATCTCT GTTGGAGCAA CAGAAGATCC GAATGTATCG CGTATCACTA	1920
TTTATTATTGA TGTGCTTCT CATGATGAAG TGGAGCAAAT CATCAAACAG CTCATCGTC	1980
AGATTGATGT GATTCGCATT CGAGATATTA CAGACAAGCC TCATTTGGAG CGCGAGGTGA	2040
TTTTGGTTAA GATGTCAGCG CCAGCTGAGA AGAGAGCTGA GATTTAGCG ATTATTCAAC	2100
CTTCCGTGC AACAGTACTA GACCTAGGCC CAAGCTCGAT TACCATTCAG ATGACGGAA	2160
ATGCAGAAAA GAGCGAAGCC CTATTGCGAG TCATTCGCC ATACGGTATT CGCAATATTG	2220
CTCGAACGGG TGCAACTGGA TTTACCCGCG ATTAAAAATC CAACTTAAAT TTATTAAACC	2280
AGCCTAAAAG GCAATAATAA ATAGAAAAGA GAGAAAAGCT ATGACAGTTC AAATGGAATA	2340
TGAAAAAGAT GTAAAGTAG CACCACTTGA CGGTAAAAAA ATGCCCGTTA TCGGTTATGG	2400
TTCACAAAGG CATGCGCATG CTCAAAACTT GCGTGATTCA GGTGCGTACG TTATTATCGG	2460
TGTACGTCCA GGTAAATCTT TTGATAAACC AAAAGAAGAT GGATTTGATA CTTACACAGT	2520
ACCAGAACCT ACTAAGTTGG CTGATGTTAT CATGATCTTGC GCGCCAGACG AAATTCAACA	2580
AGAATTGTC GAAACAGAAA TCCCTCCAAA CTTGGAAGCT GGAAACGCGAG TTGGATTTGC	2640
CCATGGTTTC AACATCCACT TTGAATTAT CAAAGTTCCCT GCGGATGCTAG ATGCTTCAAT	2700
GTGTGCTCCT AAAGGACCAAG GACACTTGGT ACGTCGTACT TACGAAGAAG GATTTGGTGT	2760
TCCAGCTCTT TATGCACTAT ACCAAGATGC AACAGGAAAT GCTAAAACA TTGCTATGGA	2820
CTGGTGTAAA GGTGTTGGAG CGGCTCGTGT AGGTCTTCTT GAAACAACTT ACAAGAAAGA	2880
AACTGAAGAA GATTGTTTG GTGAAACAAGC TGTACTTTGT GGTGGTTTGA CTGCCCTTAT	2940
CGAACGCAGGT TTGAAACTCT TGACAGAAGC AGGTTACGCT CCAGAATTGG CTTACTTTGA	3000
AGTTCTTCAC GAAATGAAAT TGATCGTTGA CTTGATCTAC GAAGGTGGAT TCAAGAAAAT	3060
GCGTCAATCT ATTTCAAAACA CTCCTGAATA CGGTGACTAT GTATCAGGTC CACGTGTAAT	3120
CACTGAACAA GTTAAAGAAA ATATGAAGGC TGTCTTGGCA GACATCCAAA ATGGTAAATT	3180
TGCAAATGAC TTTGAAATG ACTATAAAGC TGGACGTCCA AAATTGACTG CTTACCGTGA	3240
ACAAGCAGCT AACCTTGAAA TTGAAAAAGT TGGTGCAGAA TTGCGTAAAG CAATGCCATT	3300
CGTTGGTAAA AACGACGATG ATGCATTCAA AATCTATAAC TAATTAGAAA TATATAGGCC	3360
TGGAGATGAT TTTATGAAAA AGATTATGAG AAAAATTGCA TCGTTATTAT TGGTTCTAGT	3420

208	
TGTATAATGT AATTACACCG TCGGTAATAG TGCTAGCAGA CCAAAATAAA GCAGATTGGT	3480
CGTATGATGA AAATGCTGTA ATTAACATTT ATGATGATGC TAATTTGAA GATGGTAGGT	3540
TGCATATGAA CTTGAACAA TTCTTCAAT TGGCACAAAT AGCTAGAGAA GAAGGTCTG	3600
AAATTCAATTC TCCGTTGAG AGAGCTGGTG CGACTAAATC TGCTCGTTAT ATAGCGAAAT	3660
GGATTTGAG AAATAAAAAA CATTAACAAA TATAGTTGGT AAATCATTAG GACCTAAATC	3720
AGCTGTTAGA TTCCGAGAAG CTCTTACCTA TATTGAAGGT CCTCTTCGCA GAATAATGA	3780
GACGATAGAT GGCGGTTAT ATCAAATAGA GCAAATTATT GCATCTGGAT TGAAAGAAC	3840
GGGTTAAAT GACTGGACTG CGAAAACCTT AGCTTCAGCT ATTCTGGGA TATTAGATGT	3900
ACTTATTTAG GGGTGAAAT CATATGAATA TTACCAATTG GTTTCTATC AAGACAGGAT	3960
GTGATGAAAC TGATAGGCAA CTGCAAAAC TATTTTTCA GTTGGATTTA CAATTGGAG	4020
AATTGACAGA TCAACTAAGA AAATTAGATT CTAATTTGT TCCTCGTAGT CAATTTGTAG	4080
ACACGTTGGA TTTGAATGAT GTAGAAATATA AAGAAATTTT AAACATTTT ATCTTCATC	4140
GTAATGATAG TGAAGAAAGT TTGGTAGAAT GGTTATATGA TTGGATTTCC ACAATCGTT	4200
ATGAACTTCC TAAAGAGTTT TCGATTGTA TGGCTCATAA ATACCATGAA AGTGTACTG	4260
AAGTTTCGG AGATGAATAA CTAAAAAACA GTCATTAGTG ACTGTTTTT ATAGAAAAAG	4320
AGGTTTATA TGTTAAGTTC AAAAGATATA ATCAAGGCTC ACAAGGTCTT GAACGGTGTG	4380
GTTGTGAATA CTCCACTGGA TTACGATCAT TATTTATCGG AGAAGTATGG TGCTAAGATT	4440
TATTTGAAAA AAGAAAATGC CCAGCGTGTGTT CGCTCCTTTA AAATCGTGG TGCTTATTAT	4500
GCCATTTCCC AGCTCAGCAA GGAAGAACGT GAACGTGGGG TAGTCTGCCG TTCTGCCGGA	4560
AATCATGCCG AGGGAGTAGC CTATACCTGT AATGAAATGA AAATCCTGC TACTATCTT	4620
ATGCCCATTA CTACGCCACA ACAAAAGATT GGTCAGGTTG GCTTTTTGG TGGGATTTT	4680
GTAACTATTA AACTAGTTGG AGATACCTTT GATGCCCTCAG CCAAAGCAGC TCAAGAATT	4740
ACAGTCTCTG AAAATCGTAC CTATTATTGAT CCTTTGATG ATGCTCATGT TCAAGCAGGT	4800
CAAGGAACAG TTGCTTATGA GATTTAGAA GAAGCTCGAA AAGAATCGAT TGATTTGAT	4860
GCTGTCTTGG TTCCCTGTGG TGGTGGCGGT CTCATTGCCG GGGTTCTAC CTATATCAAG	4920
GAAACAAGTC CAGAGATTGA GGTATCGGA GTAGAGGCAGA ATGGAGCGCG TTCCATGAAA	4980
GCTGCCCTTG AGGCTGGAGG TCCAGTAAAA CTCAAGGAAA TTGATAAATT TGCTGATGG	5040
ATTGCTGTGC AAAAGGTAGG TCAGTTGACC TATGAAGCAA CTCGTCAACA TATTAAAATC	5100
TTGGTAGGTG TCGATGAGGG ATTGATTTCT GAAACCTTGA TTGACCTTTA CTCTAAGCAA	5160
GGGATAGTCG CAGAACCTGC TGGAGCGGCT AGTATCGCCT CTTTAGAGGT TTTAGCTGAA	5220

209

TATATTAAGG GGAAAACCAT TTGTTGTATC ATTTCTGGAG GAAATAATGA TATCAACCGT	5280
ATGCCAGAAA TGGAAGAGCG TGCCCTGATT TATGATGGTA TCAAACATTA CTTTGTGGTC	5340
AATTTCCAC AACGTCAGG AGCTTGCGT GAGTTGTAA ATGATATCCT GGGGCCAAAT	5400
GATGATATCA CACGTTTGA GTATATCAA CGAGCTAGCA AGGAAACAGG CCCAGTATTA	5460
ATTGGGATCG CTTTAGCAGA TAAGCATGAT TATGCAGGTT TGATTCGTAG AATGGAAGGT	5520
TTTGATCCAG CTTATATTAA CTTAAATGGT AATGAAACGC TTTATAATAT GCTTGTCTGA	5580
GGACTAATAA AAAAATATCA TACCTTCATT TTGATTCCT ATCTATTGAC AAGCATAGTC	5640
ACACTGTCCT TAATACTCTT CGAAAATCTC TTCAAACAC GTTACGCTCTA TCTGCAACCT	5700
CAAAACAGTG TTTTGGCAA CTTGGGCTA GCTTCCTAGT TTGCTCTTG ATTTCATTG	5760
AGTATAAGGT ATGATTTGAT TTCTTTTG TGACAAATAT ACTATATTAA AAAGATATAT	5820
AAGTAATTAA CTGAGCTTAT CTGCTTGTC ATCTCTATTAA AGGATGGTTT AGATAATCGG	5880
GTGTCGCTT CTAGGCTAGC ACCTCAATAT CCAAAGGAGT GATGAATTG AAGGACATAA	5940
GGAATACCTA TCTCTCAGAT GATTATTGAGA GGAAGAAAAGA TAGGAGTTTT TGAGCTAGTG	6000
AAGGCTTGGA TTTCTAAAGG TTAGAACTAT CATCTTCAGT TCTTAAATCG AAGAAATAAG	6060
CTATCTTACG GAAATAGAGA AGCATTTCAGT AAGAACTTGA ATAATTTCGC ACCTTAAGAG	6120
CGTAATAATA CAGTATTTT ATTAGCAAAT ATTATGGTG TAGAGGCTAG CAAAACCTAT	6180
ATATTATCGG ATTTAAAAG GAAGTAAGAA A	6211

(2) INFORMATION FOR SEQ ID NO: 9:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 7939 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:

CCGGACTCCC CACGATTCTT CAAAATAACT GACTATATT CTATCTTGAT TTTCAGATAT	60
AAATTCTTCC TTCTGTGGCC TCTTCTTACG CTTGAGAAGA GCTTCTCCGA CATGGCTTCT	120
TCCTTACTGA GCAAAACCTT GACCATAGAT AAGTTGACT GGCAAGCGTG CTCTTGATA	180
TTTGGCTCCC TTCCCACTAT TGTGGATAGC GAGGCGTCTT CTCATATCG TCGTATAGCC	240
TATATAGTAG GATCCATCAC GACACTCCAG AACGTACATA TAAGCCTTAT GATCCATAAT	300
AAATCTCTTC GATTTCGGGC GTATAAGAGC CATCATCATT GTGGACAATC AAAGGAGGTA	360

210						
AGACCTTAAA	GCCACTGTT	GAGCCATCCT	TGATCGCCTC	AATCAAAAGC	ATATTGGCTT	420
CCTTTCTCT	TTTGGATAA	ACAAACTGCA	GGCGCTTAGG	GGCTAGATTA	TGTCGTTTA	480
ACGTATCCAA	AATATCCAGA	AGTCGATCAG	GACGATGAAC	CATGGCCAAA	CGCCCATTAG	540
ACTTGAGAAT	ACTCTGGCA	CTACGACAGA	TTCTTCCAA	ATTAGTCGTG	ATTTCGTGTG	600
GAGCCAAGAG	ATAATGTTCA	CTCTCGTTCA	GATTAGAATA	AGGATTCAAC	TTGAAATAGG	660
GTGGATTACA	CAAAATCAT	TCCACCTAC	TCCCCTGAAT	GTGAGCAGGC	ATATTTTCA	720
AATCATCGCA	GATGACCTGC	ATTGCTCCT	CTAATCCATT	CAAACGGACA	GAGCGTTAG	780
CCATATCCGC	CAAACGCTCC	TGAATCTCAA	CAGACAATAT	CTGTGCTTGA	GTACGAGTGC	840
TAGCAAAAG	CCCCACTGCT	CCATTCCCAG	CACAGAAATC	CACAATCAAC	CCCTTCTTAG	900
GAAAACGTGG	AAATCGTGAT	AAGAGAACAC	TATCCACCGA	ATAGCTAAA	ACCTCTCTAT	960
TTTGAATGAT	TTTGATATCT	GTGAAAAGA	GCTGGTTAAT	GCGCTCTCCT	GATTTAATA	1020
ATTGTTCTTC	TTCCATGGTC	CTATTATAGC	AAATTCAAT	TAACATTACA	AAAAATATAA	1080
AACTCTAAC	TACTCTTCT	TTTTAAATG	GTGCAGGGCT	TCTCCAGTCC	AGATTGGTAG	1140
CATTGTCGA	AAGGGAGCAA	AGCCGTAGTT	AAAGCGGTG	CTTGAAAAGC	GTCTCCGTCT	1200
AGGAAACTGG	TACTTTCTT	CCTCCAAAGT	GGGGATAGAA	AGACTGGCTT	TCCCTGTAAA	1260
TTCATCTAAA	TCCACTACCT	GAACCTGAAC	CTCTTCATCG	ACTTTCAAGG	TTTCATGAAT	1320
ATTTTCAATA	AATCCGTGTC	GAATCTCTGA	AATGTGAATC	AGCCCCGTAT	CACCCGTCTC	1380
TAACCTAAC	AAGGCACCGT	AGGGCTGAAT	CCCTGTAATA	CGCCCCTTA	GCTTATCACC	1440
GATTTTCATC	TTAGTCCTCG	ATTCAATTAG	TTCAATTAC	AACATCTTC	ACTGGCTTGT	1500
CCATAGCTCC	TGTCTCAACA	GCAGCAATGG	CATCCAAGAC	AGCGTAAGAT	GCTTCATCAG	1560
CTAACTGACC	AAAAACCGTG	TGACGGCGGT	CTAGGTGAGG	TGTCCCACCT	TGATTGGCAT	1620
AGATTCTGC	AATCGGTTCT	GGCCAACCCAC	CACGAGTAAT	TTCTTCTTA	GAATAAGGTA	1680
GGTGTGGTT	TTGCACGATA	AAGAACTGGC	TGCCGTTGGT	ATTTGGACCA	GCATTTGCCA	1740
TGGAAAGAGC	ACCACGGATA	TTGTAAAGCT	CTTCTGAGAA	TTCATCCTCA	AAAGATTCCG	1800
CGTAGATTGA	CTGCCACCC	ATACCAGTTC	CAGTTGGTC	TCCACCTTG	ATCATAAAAGT	1860
CCTTGATAAT	ACGGTGGAAA	ATGACACCAT	CATAGTAGCC	ATCTTTGAA	AGAGATAACAA	1920
AGTTAGCCAC	TGTTTTAGGA	GCATGTCAG	GGAAAAGCTT	GATACGTAAG	TCTCCGTGAT	1980
TGGTCTTAAT	AGTCGCAAGA	GGACCTCTA	CTGTTCAAT	GTCTACTTGT	GGAAAATGCA	2040
ATTCTTTTTC	TACCATACCA	AATACTCTA	AGGCAGCAA	AATGCCATCT	TCTTCTAATG	2100
TTTTGTAAAT	ATAATCTGCT	TTTCTTTGA	TTTTATCATG	AGAAATTCCC	ATGGCAACGC	2160

TGATTCCAGC ATAATCAAAG AGTTCCAAGT CGTTGAGACC ATCTCCAAA ACCATGACCT	2220
TCTCTGGTTT CAAGCCAAGG TGTTCCACAA CCTTTTCCAC CCCCCTCGCT TTGGAGCCTG	2280
AAATCCGCAC AATATCAGAC GAATGTTGAT GCCAACGAAC CATGCGAAGT TTGTCAGAGA	2340
GAATGTCAGG CAAGTGCAAG TCATCTCCCT TATCTTCAAA AGTCCACATC TGATAGATAT	2400
CTTCTTTTC ATGGAATCG GGATCTACAT CTAAGTCGGG ATAAATTGGA TTGATAGCTT	2460
CACTCATCAT ATCGGTGCGA GTCGACAAC TGGCATCATG ACTCCCAACC AAGCCATACT	2520
CAATTCCCTTC TTGCTTAGCC CAAGAGATAT ACTCCTCAAC ATCTGACTTT TCAATCTGAT	2580
GCTGATAAT GACCTGACCT TTTTATCTT CGATATAAGC CCCATTCAAA GTTACAAAAA	2640
AGTCAGGCTT GAGATCAGCA ATCTCTGGAA CAACACCAAA AATGCCACGT CCAGAGGCCA	2700
TTCCTGTTAA AATTCCCTTT TCACGCAACT GTTTAAAAC AGTGGGAATT GTAGTTGGAA	2760
TAAACCTGT CTTTGAATTC CCAGATGTAT CATCAATATC AAAAAAGACA ATCTTGATCT	2820
TCTTGCCTT GTATCTTAAT TTGCGGTCCA TCTCACTACC TCTTCAATC TAACTCTTTC	2880
CATTATATCA TAAAGTAGGC AAATCCCCTA TTTTCAAAA GTTTATCATT TTTATTTTAA	2940
TTTCTGGAT GAGAAAAGAG ACATATTTAT GAAAAAGCTC CATCGTGCTT TTAATGTGTT	3000
CTCTTGTGTTT CAAACTCGTA AAAAGGGAGC CACTGATCCT AACTCGCTCT CTCATTTCAA	3060
AGCTTGTGAA AAAAGACCG TTGGGGCTTT AATTGCTTT CTTGTTTCA AGCTCATGAA	3120
AAAGAGACCC AACTGGGTCT TTTCTTTAAT CTTGTTTAC CAAACCCATC AAACCCATTA	3180
CGCGGACGCG TTTGATAGCT GTTGTGTTACTT TACGTTGGTT TTTAGCTGAA GTTCCTGTTA	3240
CACGACGAGG AAGGATTTTC CCACGTTCTG AAACGAAACG GCTAAGAAGC TCAGTATCTT	3300
TGTAATCAAC ATATTCAATT TTGTTGCTG CGATGTAATC AACTTTTTA CGGCCTTGA	3360
ATCCGCCACG ACGTGTTGTA GCCATGTTT TTCTCCTTTA TAAGTTTAGT TGTCCATTAG	3420
AATGGTAAAT CATCATCTGA AATATCCAAT GGGTTGTTG CTCCAAATGG ATTTTCATTA	3480
CGTGAAAAGT CTGGTACTGA ATTTGTAGGT GCTGAATAGT TTGCAAGTTGG TGCAGAGTAA	3540
GCTCCACCTG TGTGACCTC ACACACACTA CGGCTTCCA ACATTTGAA ATTCTCAGCC	3600
ACGACCTCTG TCACGTGAGAC ACGGTGTCCCT TGCTGGTTAT CGTAACACTAG AGTCTGGATA	3660
CGACCTGTCA CCCCCATAAG TGAGCCTTTT TTAGCCCAGT TAGCAAGATT TTCAGCCTGT	3720
TGGGCCACACA TAACGACATT GATAAAATCA GCCTCACGTT CACCATTTG ACTCTTAAAT	3780
GTACGGTTA CTGCAAGAGT AAAAGTCGCA ACTGCTACAT TTGATGGGGT ATAACGCAAC	3840
TCAGCGTCAC GTGTCATACG CCCTACAAAGT ACAACATTGT TAATCATAGT TTACCTTCTT	3900

212

ACCGCGTCAAT	TTTGACGATC	ATGTGACGAA	GAATGTCAGC	GTTGATTTTT	GAAAGACGGT	3960
CAAACCTTTT	AAGAGCTGCA	TCGTCATTTG	CTTCAACGTT	AACGATGTGG	TAAAGTCCTT	4020
CACGGAAATC	TTGGATTTCG	TATGCAAGAC	GACGTTTTTC	CCAAAGTTTT	GATTCAACAA	4080
CAGTTGCACC	GTTGTCAGTC	AAAATAGAGT	CAAAACGTGC	TACCAAAGCG	TTTTAGCTT	4140
CTTCTTCAAT	GTTGGACGA	ATGATATAAA	GAATTCGTA	TTTAGCCATT	GATATGTTCC	4200
TCCTTTGGT	CTAATGACCC	CAAGACTTTG	CAAGGGTAA	GTGAGGTTCG	CTCACAATAA	4260
ACTATTATAC	TAGAAAAAAAT	TTTTTACGC	AAAGTAAAAC	ACTAGAATTTC	GAAAAAACGC	4320
CACATGGCG	TTTCCCTGTT	CTTATGGTTT	GATACGGTGC	AACATACGTG	GGAATGGAAT	4380
AGCTTCACGG	ATATGTTTG	TTCCGTGCTGC	GAAGGTTACC	ATACGTTCGA	TACCGATACC	4440
AAATCCTCCG	TGTGGAACTG	TACCGTATTT	ACGAAGGTCA	AGGTAGAATT	CATATTCTGT	4500
ACGATCCATG	CCAAGTTCAT	CCATCTTAGC	GACAAGGGCA	TCGTAATCTT	CCTCACCGAT	4560
AGACCCACCG	ATAATTCTC	CATAGCCTTC	TGGAGCAAGC	AACTCTGCAC	AAAGCACGCG	4620
CTCTGGATTT	CCAGGAAC TG	TTTCATGTA	GAAGGCCTTG	ATGGCTGCTG	GATAGTTCAT	4680
GACAAATGTT	GGCACACCAA	AGTGGTTGA	AATCCAAGTT	TCGTGTGGTG	ACCCAAAGTC	4740
ATCACCATGC	TCAAGATGCT	CGTAGTCAGC	ATCTTCATCA	TTTCATGCT	CTTGCAAGAG	4800
GTCAATGGCT	TGATCGTAAG	TGATACGTTT	GAATGGCTCT	GCAATGTAGC	TTTCAAGAG	4860
TTCTGTATCA	CGTTCCAAGG	TTTCCAAGGC	TTGAGGCGCG	CGGTCAAGAA	CACCTTGAG	4920
AAGAGCTTTC	ACATAAGCTT	CTTGCAAGTC	AAGCGACTCA	TCATGTGTCA	AGTATGAGTA	4980
CTCAGCATCC	ATCATCCAGA	ACTCAGTCAA	GTGACGGCGT	TTTTTGATT	TTTCAGCACCG	5040
GAAAACGTGGA	CCAAAGTCAA	AGACACGACC	AAAGGCCATA	GCCCCCTGCTT	CTAGGTAAAG	5100
CTGACCTGAT	TGGCTCAAGT	AGGCTGGCGT	TCCGAAGTAG	TCAGTTCAA	AGAGTTCTGT	5160
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GTCAAAGAAC	TCATAAGTTG	CATAGATAAT	AGCGTTACGG	ATTTGCAACA	CAGCTACTTG	5280
CTTACGAGAG	CGTAgCCACA	AGTGACGTT	ATCCATCAA	AACTCTGTTC	CGTGTCTTT	5340
TGGTGTGATT	GGGTAGTCTT	GAGATTCAACC	GATCACTTCG	ATGTCTGTGA	TGTCCAACTC	5400
ATAGCCAAAT	TTAGAACGTT	CGTCCTCTTT	GACAATACCT	GTCACATAAA	CAGACGTTTC	5460
TTGGCTCAAG	CGTTTGATAA	CATCAAACCTT	CTCAAGTCCC	ACTTCTTCAC	CAAATTTTTC	5520
GACAAAGTTT	GGTTTAAAAG	CCACACCTTG	AAAGAAGGCT	GTTCCATCAC	GCAATTGTAA	5580
GAAAGCGATT	TTTCCTTTTC	CTGATTTGTT	GSACACCCAA	GCGCCAATCG	TCACCTCCTG	5640
ACCAACATAG	TCTTTTACGT	CAATAATCGT	TACACGTTTT	GTCATTATTT	TTCCCTTTCT	5700

TTTTTATTCT TTATGGCAA CCACCTCTAT ATTGTTCCA TCCAGGTCAA TCATAAAAGC	5760
AGCATAGTAA ATCGGATGCT CACTTCGATA ACCAGGAGCC CCATTGTCTC GCCCACCTGC	5820
CTCTAAGCCA GCCTCATAAC AAGCCTGAAC TTCTTCCTTA TTTTCTGCTA AAAAAGCAA	5880
ATGAACAGGA TCTTGTGTTC CCTGAGTCAG CCAAAAATCA CCACCAGGAT GAGGGCTGTT	5940
CGGGGATAGA AAACTAATTA GAGAACTAGT CTTAAAAGCC AATTATAGT CCAAAGGAGC	6000
GAGAAAACTC CTATAAAATC CTTATGAAAT TTGTAAATCC TTTACCTTAA TCTCAAAATG	6060
ATCAATCATT CTCACTACCC ATAAAATGCTT TCAAGCGTTC GACTGCTTCT TTAAGCGTGT	6120
CTAGGTCTGT CGCATAGCTG AGGCGGACAT TTTCTGGTGC TCCAAATCCA GCTCCTGTTA	6180
CCAAGGCCAC TTCCGGCTCT TCTAAAGATAA CAGTTGTAAA GTCTGTCACA TCCGTTAGC	6240
CTTTCATCTC CATGGCCTTT TTGACATTTG GGAAGAGATA GAAGGCCCC TGCGGTTGA	6300
CCACTTCAAA TCCTGGTACC TCTGCAAGGA GGGGATAGAT GGTATTAAGA CGTTCTCAA	6360
AGGCCTGACG CATGCTTCT ACAGTATCTT GCTCACCTGA TAGACCTCA ACTGCTGCAT	6420
ATTGGGCTAC TGCTGACGGA TTCGAAGTGTG TTTGACCTGC AATCTGGAC ATGGCAGCGA	6480
TAATGCTGC TTCTCCAACG GCATAACCAA TCCGCCAACC AGTCATGGCA TAAGTTTAG	6540
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TGAACTCATG ACCATTATAA ACCAACCGGC CATAGATATC GTCTGCTAGG ATGAGAATAT	6660
CATTTCATCAG AGCCAGTTT CCAATTGCCA AGAGTTCTC ACGGGTGTAA ATCATACCTG	6720
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ACTGCTCTAC GGTCACCTTA AAGTGATTGT CTTCTCTAGC AGAACAAAG ACGGGAACGC	6840
CTTCTGCCAT CTTGACCTGA TCTCCATAGC TAAACCCAGTA TGGGGTTGGG ATGATGACTT	6900
CATCACCTGG ATTGACCCACA GCCATAAAAGA AGGTATAGAG AGAACATTTG GCTCCCGCAG	6960
CGACTGTCAC TTGATTTGAC GCTACAGAAT AGCCGTAAAA GCGCTCAAAG TAGCTATTGA	7020
CCGCCGCCTT AAGCTCTGGC AGACCTGAGG TTACTGTATA AAAAGAAGCA CGCCCATCTC	7080
GAATCGATCC AATGGCGCA TCTTGGATAT TTTTGGGAGT AGTAAATCT GGCTCACCC	7140
AGGTTAGAGA CAAAATATCT CTACCCCTCAG CCTTCAGTGC TTTGGCACGG GCTCCAGCAG	7200
CCAAAGTCAC ACTTTCTTCC ATTCTAAAA CACGGTTGGA TAGTTTCATA GGCCCTCCTT	7260
GTTGACCAAT GCTCTGTCTT CAAAATCTAC TAGATAAAA TCAGATCCTG ACTTAACTTC	7320
CCAGATTGGC TTATCTGTAT AACGGCCAAA GTTATCTTG TCAATCTCGC CAGCTCCCTT	7380
TTCCTAGAA ACCGTTCTG CTTTTCTTG TGAAACACCC TGATTTAGCT GATAAACGTA	7440

214	
AATCTTATGG TCATCTTAC CAATCAGGAC AGCAAGCGCT TCTTGCTGTT TCTTACGACC	7500
AAGAACGCTG TAATAAGATT CCAAGCCATT GTATAATCA ACCTGATCAG CCTGCTCTAA	7560
TCCTGCATAC TGCTGAGCTA ATTTTTCTCC TTCACTTTA CCTGTTTGAT AGGGTTTCAT	7620
GCTAAGAGAA ACCATATACA GAAAGGAACC ACTGATAACC ACAAAACAAA TCGTCATCCC	7680
TAGACCATAC TGCCACAGTA GATTATTTTG TGCTTTGTT TGTCTTTTT TCACTCGTCT	7740
ATTTTACCAT CTATTAAAGCT TTATTACAAG TGAATATAAG AATACTCTTC GAAAATCTCT	7800
TCAAACCCACG TCAGCTTTAT CTGCAGACCT CAAAGCTGTG CTTTGAGCAA CCAATTCTAT	7860
TTCTCCCTTC AAACAAAACC GATTTTGAAA GTGAAACAGT TCTTACTTTT TCAGTCACAA	7920
ATGATTAGAG TTTGCCGGG	7939

(2) INFORMATION FOR SEQ ID NO: 10:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 9897 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:

CCGCTCTPACC GTCAAATAAT TACCATTTG TTTAATACCG AAATTTTTAT CTACTGAAAA	60
TTCAGTTGGT CTGTTGGTAC GATCGTCGTA TACAGTACCA TTCTCACGAA TAGTATAATT	120
GTAATCAGTA TCACCTTGTT TCCTTAATTT AAGGTAATAA TTACCATCAA TTTGTTTATA	180
ACCTGAATCT TTTCTAGTTG CTTCTCTAAA ACTTACTCCA GCAGGCATCA CATCAGCAA	240
CATGAGTACT TGTTTGTCT TTTTTCAAC AATAACAGAG TCAATATAGG TTGCACCACC	300
GCTGATTGT AAGTCACGTC CACCAACTTC ACGAGGCCAT TCTAATGGTA CTGGCGCAA	360
ATCATCGAAT GCCAATGTTA ATTTGGTTT AGTCCATGTC TTACCAATTAT CATCACTATA	420
ACTTGTAGCA ATATTAATTT TATTCAAGAA ATCATGAGTT CCACCGTAAC GAGCGTCAAT	480
GCTTAAAAAT ACCCGACCAT TGCTAAAGT ATACAGAACT GGAATACGGA AATAGTTAGA	540
ACCTGTGTA TCATTTAGCCG TATAAATTAA ATGTCCAGTA ACAGCGTTG TTGTCATCTT	600
TTTAACAGTT TCTTCATCCA ATGCACTATT AAAGAATTG ATATTTCTA GTGTTCCGTT	660
AAAACCAAAAC GCCGTTTTTC CTGCACGTTT CACTCCCCA AGCATATACT AATCAATACC	720
TTTAATATCC TTGATGTTTA GGAAATTATC CACTTTCTTT TCTACTACTT TTGTACCAATT	780
TGCGTATAAA GAATATGTTT TTTTGACTGA ATCTGCTACT ACTGCAACAG TGTTAGTCAC	840
AGCCTCTTGT TTGTAATTAC CCCAAACTGA AGCAGGTCTG GATACTAGGT TATTTTTATT	900

GGAAGAAGTA	TCACCGCGCTT	CCATCCCCAA	CTCACCAATTG	TCTCTAAGGA	ACACATCTAC	960
ATAACTATTT	TGTTGACCGG	GTTTGGATT	AGATATTCCA	AACAGAGCTT	GTAAGCCTT	1020
CTCACCTGAC	TGATTGTA	TAATCACTAC	AGTAAAGTCA	CCGCTAGTAA	ATTTATCCTT	1080
TAACCTTTA	GTAACATTTT	CTCCGCCCCC	TGTTAAAGTA	ACATTATTTT	TTTCTAAGAC	1140
AGGAGTTCT	TCCGCTGTAG	AAGATGGATC	CTAACAGTA	GTTTCAACTG	TTCGAGGTTG	1200
TACAGTAACT	TCCGAAGAGT	TATCCGATGT	AGGTTGTA	TCCGAAATCG	GAGTCGTTGG	1260
TGCAACAGGT	TGCACCAACT	TTGGTGTGAA	TACTTCAGAA	GTTTCAGTCT	CCTGAGCTGC	1320
AACTGAGTTA	GCAACAAATG	CTGATAATAC	CACTACAGTA	CCTAACAGTTA	CATATTGTTT	1380
AATATTTTTT	TTCATTTTAT	TTTCCTCGT	TTAAACATT	GATAACAACT	TTTTAACAG	1440
TTTCATCATT	GCAATGAATC	TTGGTTGGT	GAAGATCTTC	TTCAAAAGTC	ACCAACATAT	1500
TCCCTGGAAG	CAATTCAACA	ATTTGATAGT	CTTTGCTATC	GTAAAAAGCA	ATATCCTTCT	1560
CTTCGCTAAA	AGGTACACGT	GACTGGGCAC	GAACCTGGGA	AGTTACTGCC	ATTTTTTCAG	1620
TATTTTCAAC	AAACATATGA	ATATCTAAAT	ATTTCCTATG	AGTTTCAAAA	ATATCTCCTG	1680
GAACCTCCATC	AGCTAGATAA	GTCATACAAT	TTGCAAAAC	ATTTTCCCCG	TCAATATCAA	1740
TTTTTCCATC	AACTAAATCT	GTCAAATTG	TATTTCTAA	AAAATCACAG	ACTTTTGAAA	1800
AATATTTATT	GACAGAAGCA	TATCGTTAA	AATCAGATTG	TTCAGAAATA	ATCATATTAT	1860
TTTCTCTTTT	CTATTAGTGA	CGAACTTCCC	AACTTGAATC	CGCTTTAATT	TCTGTAATAT	1920
CATGAATCGT	TGTATATTAA	GGTGCAGATA	CTTATTTCC	AGTAAGAAC	GATACAATAT	1980
AACCTGAAAC	TACTGATACA	GAGATTGAAA	TCAATGAATA	TGCCCACTAG	CTAACAGCTG	2040
TTGGAGGAAG	GAAGTATTAA	ATAAAATCCA	TGACGATGGT	TGATACAATC	AGCGCTGCAT	2100
AAGCACCTG	TTTATTTGCT	TTTTTAGAAA	CAAATCCAAG	AATAAAATACA	CCACCAAGTA	2160
GACCAAGTAC	AAGTCCCAG	AAACTATTGA	ACCATTGCA	TGCAGATTAA	ATATCTGAGT	2220
GAGCCATGAC	AATGGAAACA	CCAATTGAGA	ATAAACCTAC	TGCTAGAGAT	ACGAATTGTG	2280
CAATTTTCGT	ACGACGATTG	TCTGACATAT	TTTTAGAAAT	GACATCTTGA	ATATCCAATG	2340
TCCATGAAGT	TGCAACAGAG	TTCAAACCTG	TTGAAATAGT	TGATTGAGAT	GCTGCATAAA	2400
TCGCTGCCAA	GATCAAACCT	GTGATACCTA	CTGGTAACTG	GTATGCAATA	AAAGTACATAA	2460
AGATTTGGTC	TTGAGGGATA	TTGCTAGCTG	CACTATCTGC	ATTTTGTACT	TGATAGAATA	2520
CGTACAAGCC	TGTACCAATC	AAGTAAAAGA	CTGTTGCAGT	TGCAAGTGAC	AAAACACCGT	2580
TTGTGAACAA	CATCTTATTA	AGTTTCTTAA	TATTTTGTGT	TGTAGTAAAA	CGTTGAACCA	2640

216

AATCTTGAGA	TGAAGCATAG	GAAGACAAGA	TTGTAAAGCC	TGAACCCATC	ACAATTAAAA	2700
AGATGGAGTT	TGAAAGCAAG	TTAGGATCGA	AAAGTTTTTC	ATTTGCAGCA	AGGAATTCC	2760
CGTTTGTCAA	TGTTTCTGCT	ACTGCACCAA	AGCCACCTTT	AATATTAGCA	ATCAGTACAA	2820
ATAAAGCTAA	AACCACACCA	CTAATCAGAA	TCACACCTTG	AATAAAGTCT	GTCCATAATA	2880
CGGATTTAG	ACCACCAAGTA	TAAGAATAAA	CAATTGCAAC	TACACCCATC	AAAATAATCA	2940
AAATATTGAT	GTCAATTCCCT	GTCAATTACTG	ATAAACCAGC	TGATGGGAGG	TACATAATGA	3000
TAGACATACG	TCCCAATTGA	AAATAATAAA	ACAAGAGTGC	TGAAATAATA	CGAAGTGCTT	3060
TAGAATTAAA	ACGTTTATCC	AAAGTAATCAT	ATGCCGTATC	GATGTCTATC	CGTGCAAAGA	3120
TAGGTAAGAT	AAAACGAATT	GTCAGTGGAA	TAGCTACTAC	CATCCCTAAT	TGAGCAAACC	3180
ATAAAATCCA	GCTACCTGCA	AAAGAGCTAC	CAGCGAGTCC	CAAGAAGGAA	ATCGGACTGA	3240
GCATTGTGGC	AAAATGGAT	ACCGAAGTAA	CATACCAAGG	AAACCGAACCA	TCTCCTTTAA	3300
AGAACTCTTT	TCCTTTCATC	TCTTTTTAG	AGAAATAGAT	ACCTGCAACC	AAACCCGCAA	3360
GTAAATAAAC	AATCAAGATA	ATTAAGTCAA	TTATTGTAAA	TCCTGTTGTG	CCCATAACAT	3420
ATCTCCATAT	TGATTTTATT	TATTATAAAA	ATTCTTTTCG	TGCTTGTGAA	ATAAGTTCTG	3480
CTGCTGTGTT	TGCAACTTCC	AAAGTCACCTT	CTGCCAATGC	TTCTAAAGGT	TGACGAACAG	3540
AACCTAAATC	AAGTTTTCA	TTTAGACGCA	AAACTCTTT	TTGCTACAGCA	TACATATTTG	3600
CCTTACCTGA	TATCATCTTA	TAGATAACTT	CATTGATAGC	ATATTGAAGT	TTTTAGCTG	3660
TATCTAAATC	TCGTTCTGTA	ATCAAACCTT	CCAATTCTAA	GAACAAATCT	GGCATAACCG	3720
CATAAGTACC	ACCAATACCA	GCTCTGCTC	CCATCAAGCG	ACCACCAAGA	TATTGTTCAT	3780
CTGGACCATT	GAATACAATG	TAATCTCTC	CACCTGCAGC	TACAAACATT	TGAATATCTT	3840
GTACAGGCAT	AGAAGAATT	TTAACTCCAA	TCACACGAGG	ATTTTGACGC	ATTGTTGCAT	3900
ACAAACTACC	AGTCAACGCA	ACCCCTGCCA	ATTGTGGAAAT	ATTATAGATA	ATAAAATCTG	3960
TATTTGACGC	AGCTTCACTC	ATTGCATTCC	AAATATGCTGC	GATTGAATAC	TCTGGCAATT	4020
TGAAATAAT	AGGTGGGATA	GCTGCAATAG	CATCGACTCC	AAACACTTTCT	GAATGTTTGT	4080
CCAATTGAT	ACTATCTTTC	GTGTTATTAC	ATGCAATATG	GTTGATAACT	GTTAATTTC	4140
CTTTAGCAAC	TTCCATAACA	GCTTCATAAA	TTTGGTTACG	ATCTTCTACA	CTTTGGTAAA	4200
TACATTCACC	TGAAGAACCA	TTTACATAGA	TACCTTTAC	ACCTTGTCA	ATGAAATATT	4260
GTACCAAGAGA	TTTACACGA	TCTGGCTAA	TTTCACCATT	TTCATCATAG	CAAGCATAAA	4320
ATGCAGGGAT	AACGCCCTTG	TATTTAGTTA	AATCTTCAT	CAGATTCTC	CTTTATATTG	4380
TTTTTATTT	GATGACATTA	ATAAATCGCT	GAGCAATTTC	TTTGGACGT	GTAATCGCTC	4440

CACCAATGAC TACACTGGTA ACACCTAAC TATAAGCTTT TTTTAATTGT TCTGGATAAT	4500
GAATTTTCT CGGCAATTA CCGGAATATT AAAATCAGCC AATTTTTCA TTAGTTCAA	4560
ATCAGGCTCA TCTGATTGTA CACTTGTACT TGTGTAACCT GATAATGTT TACCAACAAA	4620
ATCAACGCCT GATTTAAATG CATAGAGACC TTCATCTAAA TTACTTACAT CCGCCATCAG	4680
CAATTGATTC GGATATTTT CTTTTATTT TTTGATAAAAT TCAGTACAA CTAAGCCATC	4740
ATATCTTGGT CTTAAAGTTG CATCAAATGC AATGACTGTT GTTCCGCATT CTACAAGTTC	4800
ATCTACTTCT TTTCATCGTAG CAGTAATATA TGGTTCTGA GGTGGATAAT CCCTTTGAT	4860
AATTCCAATT ATGGTAAAT CTACTACTTT CTGAATTGCT TTAATATCAC GCACAGAATT	4920
TGCGCGAATG CCCACTGCTC CTGCCCTCAA AGCTGCTTTA GCCATAAAAG GCATCAAGCT	4980
AAATTCTTCA TTATAAAGGG CTMCACCAGG TAAAGCTTGA CAAGAAACAA TGACTCCACC	5040
TTGAACTTGG CTTATAAATT TTTCTTTAGT CCAAATTGG CTCATTAT TATTCCCTCCT	5100
TATGGATAAT AGTTTGATTG TAATAATATT GTCTCTCTGG ACTTTCCAGA TAATTAGAGA	5160
ATAAGCAGTC TGTAATTAAA AGTATTGGAA ACTGAGGTGA TATGCGATTG CCATACGAGA	5220
GATGATCGGT CGAAGCTAAT AACAAATAGTT CATCAAAGAA ACAATCTTCT TCGTCAAATT	5280
TTCTTGTAGT CATTAAAATC GTTTTAGCGC CTTTATCTGC AGCTTTTG AGACCTTCTA	5340
GTACAATATC AGTTTGACCT GAAATGGATG CTCCAATGAC AAGGCAATT TCATTAAGTA	5400
GTAAAGCTACT CCACAAAATC ATATCCTCGT CTGATAATAC TTCACCAATC ACTCCGAGAC	5460
GCATAAAATCT CATCTTCATT TCTTGTAAG CAAGAACAGA ACTTCCTTTA CCGTAGAGAT	5520
ATACACGCTC AGCAGTTCT ATCATCTCG CAATACGCTC AAGTTGAAC TCAATCAAGAA	5580
CCGTGTAAGT TTTCTCAAC ATTTCCCTCAT AGTCGGATAA AACTTTTCT GTTGCCTCTG	5640
TATATAATGC CAACTTTCT TTCTCATGAA TCATCTCTTG GTATTTGAAA ATGAATTGTC	5700
TAAAACCTTT AAAACCACAT TTTTCGCAA ATCGAGTCAA TGTTGTTTG GATACATTAA	5760
GGTATTGCA CAATGCTTTA GATGAATAAT CATTAGAGG TTGCTGTTT AAGAAGAATT	5820
TAGCAATGTC TTTTCAGCA TATGCCATAT TTGGTAAGTT AGCTTCTATC ATTGGAATTAA	5880
GTTCTTTTG CAGTAACATA TGAGCTCCTT AGTTGAAGTA AACGTTTACA TTCTTTATTT	5940
TAACACTTT TTTTTTTTC AATATTTTC ATAATTAGA AACTAGTTTC CAATTCTTT	6000
CGTTTCATAA CAGAACACAA AACATAAAAA TATAATAGT TTTATTCTTT TTATCGTAAT	6060
TATATGTATT GTAAGAACGT TTATCACTAA TAATATGTTCA ATATTAATAT ATTGTTAGTAA	6120
TATTTTATTT TGTTTTTATT ATTTCTTTTC GGAATTCTA TATAATATTT TATTTCTAAA	6180

218

AAAATTGAAA	AAATATTCT	AGTTTCTTTA	TTTTATATAG	GTAATATATT	TTATTTCTAA	6240
ATTAAAAGAG	AATCCCATAA	AAACTACAGA	TTTATGAGAT	AAATCAGGTC	ACCTATTTA	6300
AAAAGCAGC	AAACTATAAA	CTAAAAAGTT	CCACACCCAA	TGTAACCCCA	TACTTCCCA	6360
TAAGTCAGAT	TTATAGCGCA	CCATACCTAA	AAACATTCCA	AGTGAACGT	ACAGACACCA	6420
AGCTAGAATG	GTTCTGGAT	GATGTACTAA	GGCAAATAAA	ACACTTGTCA	AAGCAACTCG	6480
AATATCTAAT	TTTCTAACCA	AGTTCATCAA	ATTTCACGA	TACAGAAATT	CTTCACCAT	6540
ACTCGCATTG	ATTAAGAACAA	ATAAAAATGA	AAACCAAGGA	ACTTGATGTT	GAAGGCCAAT	6600
TAAATTGTT	TGATTCGTGC	TTCTTGAGC	ATGAATCAGG	CTAAACATA	GACTTATAAT	6660
CAGTAGACTA	GCTAGTCCAA	TACCAAGGCA	TTTCATCCTA	GTTTTCATAT	TGACCTTGAC	6720
CACTTGTGTT	CGTTGACCAT	ACATCCATAA	AAAAGAAAAA	AGAGACGCAC	CATAGAGAAC	6780
CTGTTAGTATA	GTAACTCAC	CGATAACAAAG	AAATTCAAT	AAAGTATAGAG	ATACCAATAG	6840
GACATTACT	TGTTGAAATA	TATAAACTGG	ATTATTCTT	TTCATAGTTA	CCTCCGAAAT	6900
AAATCTTCAT	AATCTAAATC	TAATATCTGC	ACAATCCTT	CTACCCATGG	ACTTTGAGGC	6960
ATTCGTTGTT	CCATCTTGTA	GTGGCGAATC	TTTTGATATA	AAACGATTCAA	TTCACTTGGA	7020
TAGTGAAACT	CTCCCGAAA	CATTTTCTG	GTAACTCAA	TCCAGCTGAT	ATTTCTTCA	7080
GCCAAAATAA	TGGACAAGTT	CTCCCAAAT	CGTTCAGCCA	TATTCTTCT	CCTTTAGTTA	7140
GATAAAATAA	GTGTTGyGC	CATGTAATC	AATTGTTTCG	TATCTCTTG	CAATAGAGCT	7200
CTAGCCTCTT	CCAAATTCAAG	ACTGGATAA	ACCCGCTTAT	TTGAAACAC	AAAAGGAAGT	7260
CCGATGGTTA	GTTCAAGGATT	TTTAAAATT	ATCTAACGA	AATCCGTTAA	TCTTAGATG	7320
TCACGGTTCT	AAATACGTAA	AAATGGGA	GATAAAAACT	CAAACAAATC	TGAAGAATAG	7380
CTCATCATCT	CAATTAATTT	GTCTTTGTC	ATTTCAGAAA	CTGAATGACA	AGATAACCTCA	7440
ATGCCATAGT	TTTGGAAAGAA	GTCTAAAAGA	AGTTGATTTC	TTTGGCTATT	TTTACTTAGA	7500
TAGAGATCAA	TCATGGGAGA	CCTCCAACAA	ATTTGCTTCC	ATTTGATATT	CTGAGACGAT	7560
TAAGGAATCT	AAACAACCTTG	AGAAGTTAAT	CGATTCTTG	TCTTCATCAT	AAGCTTTAC	7620
AGTTACTTGG	GTTGTAAGTA	TCCCCTCTT	TCCCTCGGCT	CGATAGTCTT	GTCATATAAA	7680
AAACAAAACA	AGATTCTGAT	TATCATCTAC	AAAGGCATTA	ACTCCGTTCT	TTATATCCTG	7740
ACTTTCAAGG	AATTCCATAA	CGTTTGAAG	ATAGGATTCA	AAAAATAGTG	GGTAATTATG	7800
TTTTTTATGG	TAATCATCTA	AAAATGTTAC	CTCAAACCTCA	CATGGATAAT	TGGGCATCAA	7860
AAATATTGTT	TCATCCAGCT	GTTGATTTTC	TGCATCATGT	AATTCTGTTT	CTAATTCATC	7920
ACAATCTAGT	ATTGATTCTT	TATTAATGC	TTTTATCTTT	TTCCCTCTATT	TCTTTAAATT	7980

TCTTTGCGAT	TGCGGCAATC	ACAGGAACGG	TTACACTATT	ACCAACTTGT	TTATAGAGCT	8040
GACTATTAAT	AGAGACTTTT	CTAGCAGCTT	CAAAGCCTA	ATCAGGAAAG	CCATGCAATC	8100
GAAAACACTC	TTTAGGAGTG	ATTCTCGTA	TTCTCAAAACG	GTAAAATTGT	CCATCTATTA	8160
AAACACCAGC	TACTTGGTAA	ACTTGTTTAT	CTTCTCCTTC	ATAGCTAGCC	ACTACTACTC	8220
CCATTGACC	ACTAGTTGTT	AACGTATTAG	CTATACCTTT	TCCAACCTCTA	CCACGACGAT	8280
ACTGAGAACT	TGGTCTTCT	AAATTGATMG	AATCCCCAAT	CTCTGCTTGA	GCATATCCTT	8340
TTTCGTTGC	TTCCCGTACT	TTTAGAAATT	GGATTGGTTC	TGGAATTAGT	ATTTGGGGA	8400
TTTTATCTCC	TCCTTGCATC	GTAGTCAGTG	TTGGAGATAA	GCCCTCACTT	CCATAGACAC	8460
GACCTGTCTC	CTTAAAGCTA	GTCGGTAAAT	CTCCAACAAAC	GACAATGCCA	TAACGATCCT	8520
GAGTATTTAA	AGTAAACATC	GGCTCTTGAT	TTTCCTTAAA	GCGTCTCCCA	TTTGTCTCT	8580
TGTCTAATCT	ATCTGGTGT	ATACAAGGAA	TCGCACTTT	AAATCCTTCT	CCTTACAC	8640
GAACTAAGGT	TGGCGCAAGA	CCTTCTGAAT	AATAGACTTT	ACCGCTCATT	CCACTTCTTG	8700
ATGGATTCAA	ATTTCTAGT	GCTTCAAAG	TCTCAGAGTT	AGTTGCTTGA	CCTTCTCGTC	8760
TGAAAGGAAA	TAAGAGTCTG	GTACCTTCT	TTCTAGAATG	TCCGATAATA	AACACCCCT	8820
CTCTGTTTTT	GGGAACGCCA	AAATCCTTAC	TGTTAACGAC	CTGCCACTCA	ACATCAAACC	8880
CCAACATCATC	AAGTGTGTA	AGTATTGTGG	TGAACGTCCG	TCCCTTATCG	TGATTGAGTA	8940
GGCCTTAAAC	ATTTCAAGA	AAAAGAAAAC	GTGGTTGGAT	TTGTTGGCC	GCCCCACCAA	9000
TTTCAAAGAA	CAAAGTTCT	CTAGTATCTT	CAAATCCCA	TCGTCTTCT	GCGATTGAAA	9060
ATGCTTGACA	AGGGAATCCC	CCACAGATGA	CATCGACTTT	CCCTCTAAGT	TTTTTAAATT	9120
CGTCATCTGA	AACATCTCGT	ATGTCATGAA	ATTCTATTT	TCCTTCCGTT	TGAAAAATGG	9180
ACTTATAAGA	TTTCCTAGCA	AATTATCAA	TCTCACAAA	TCCCAAGCAC	TCATGCCCTT	9240
GAGCTTCCAT	TCCCATCCTA	AAGCCTCCTA	TCCCAGCAA	TAATCTAA	ACCCAAATCA	9300
TTCATACCTC	TCTCAACTAG	ATGTAACCTA	CAAACCCCT	GACCTCATGA	GCCACTTTCT	9360
TCCTCCCAT	GAGGTCACTT	TTACTTTCTG	CTGTTCCAGT	ATCGTTTTTC	CTCGCTAGAT	9420
TTCCCTAAAA	GGGCAGACTC	CTCCCTTGGT	TCGTACACG	ATTTTTTCAT	CTCGACTGTT	9480
CTTTAATGCA	TCATTAACGA	CGCTTTCTT	CTAGGTGGTT	CATAAGGAAC	AGGAAGATTC	9540
AGGTTGACTT	TTCTAATCCT	AGAATAAAAGT	GCTGAAAACA	ATTCGGAATA	GGCATAGAGA	9600
CTAGACAAATT	TGAGGAGCTG	CTTGCCTCCT	GTTCGAACAC	ATTTCCCTAC	CACGTGAAGA	9660
AAAAGATGGC	GGAAAGCGTTT	GATTGTTAAA	GTTTGGAAAGT	CACCTCCAGC	TAGATGTTG	9720

220	
AGAAAAAGAT AGAGATTGTA GGCGATACAG CTCATCATCA TACGAACCTCG TTTTGATTA	9780
AGGTTGAAC T ATCCGTTTA TCGCCAAAAA ATCCCTCCTT CATCTCCTTG ATGAAATTCT	9840
CGGCTGACC ACGTCCACGA TAAAGCTGAA ACTGGTCTTG GCTTGTCCG GTACCGA	9897

(2) INFORMATION FOR SEQ ID NO: 11:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 8148 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11:

CCGTGGAACA AGCCAAGACC AGTTTCAGCT TTATCGTGG A CGTGGTCAAG CCGAGAATT	60
CATCAAGGAG ATGAAGGAGG GATTTTTGCGATAAAACG GATA GTTCAA CCTTAATCAA	120
AAACGAAGTT CGTATGATGA TGAGCTGTAT CGCCTACAA CTCTATCTTT TTCTCAAACAA	180
TCTAGCTGGA GGTGACTTCC AAACTTAAC AATCAAACGC TTCCGCCATC TTTTCTTCA	240
CGTGGTAGGA AAATGTGTC GAACAGGACG CAAGCAGCTC CTCAAATTGT CTAGTCTCTA	300
TGCCTATTCC GAATTGTTT CAGCACTTTA TTCTAGGATT AGAAAAGTCA ACCTGAATCT	360
TCCTGTTCC TATGAACAC CTAGAAGAAA AGCGTCGTTA ATGATGCATT AAAGAACAGT	420
CGAGATGAAA AAATCGTGTG ACGAACCAAG GGAGGGAGTCT GCCCTTTGA GGAAATCTAG	480
CGAGGAAAAA CGATACTGGA ACAGCAGAAA GTAAAAGTGA CCTCATGAGG AGGAAGAAAG	540
TGGCTCATGA GGTCAAGGGT TTTGTAAGTT ACATCTAGTT GAGAGAGGTA TGAATGATT	600
GGGTAAATAC AATGAGCTTG AAAGAAGTAG CAAACTCACC AAGCGCCAAT TCTTGAGAA	660
TCAGATGCTG GATTATACCA TCATTGCGCA TGAGAGTTT GAAATCATCC GTCATTCTGT	720
CTACCAGACA GATGATCGTGA AAGTGGAAAA TGCTCTGGCT TTTGAAGTGA AAAATGATGA	780
AACAGACAAG CTGATTCTGT TATTAAGCGA GGATATTGGT GTAGGTGAAA AATTGTGCCT	840
CGTTGACGGA ACAAATGCTGTTAGTAT GATAAAATAA ATGAGAGAA	900
GATTGCTTG CAGTGCTAGA AATAGGCATT TTGAATAGTG AATATGTTAT AATAAGTATT	960
AGTAGGAGGT GTTTAGATT GGAGAAGAAA CTGACCAAA AAGACATTGC GGAAATGGCT	1020
CAGACCTCGA AAACAAACCGT GTCAATTAC CTAACGGGA AATATGAAAA AATGTCCCAA	1080
GAGACACGTG AAAAGATTGA AAAAGTTATT CATGAAACAA ATTACAAACC GAGCATTGTT	1140
CGCGTAGCT TAAACTCCAA ACGAACAAAA TTAATCGGTG TTTTGATTGG TGATATTACC	1200
AACAGTTCT CAAACCAAAT TGTAAAGGGA ATTGAGGATA TCGCCAGCCA GAATGGCTAC	1260

221

CAGGTAATGA TAGGAAATAG TAATTACAGC CAAGAGAGTG AGGACCGGTA TATTGAAAGC	1320
ATGCTTCTCT TGGGAGTAGA CGGCTTTATT ATTCAAGCCGA CCTCTAATTTC CCGAAAATAT	1380
TCTCGTATCA TCGATGAGAA AAAGAAGAAA ATGGCTTTTT TTGATAGTCA GCTCTATGAA	1440
CACCGGACTA GCTGGGTTAA AACCAATAAC TATGATGCCG TTTATGACAT GACCCAGTCC	1500
TGTATCGAAA AAGGTTATGA ACATTTCTC TTGATTACAG CGGATACGAG TCGTTTGAGT	1560
ACTCGGATTG AGCGGGCAAG TGTTTTGTG GATGCTTAA CAGATGCTAA TATGCGTCAC	1620
GCCAGTCTAA CCATTGAAGA TAAGCATAACG AATTGGAAC AAATTAAGGA ATTTTTACAA	1680
AAAGAAATCG ATCCCGATGA AAAAACCTCG GTATTTATCC CTAACGTGTTG GGCCCTACCT	1740
CTAGTCTTAA CGGTATCAA AGAGTTGAAT TATAACTTGC CACAAGTTGG GTTGATTGGT	1800
TTTGACAATA CGGAGTGGAC TTGCTTTCT TCTCCAAGTG TTTCGACGCT GGTTCAGCCC	1860
TCCTTGAGG AAGGACAACA GGCTACAAAG ATTTGATTG ACCAGATTGA AGGTCGCAAT	1920
CAAGAAGAAA GGCAACAAAGT CTTGGATTGT AGTGTGAATT GGAAAGAGTC GACTTTCTAA	1980
AATGAAGGAA AATGACTTGC AATCTCTGTT AAGAAATAAA ATAATCCAC CTAGAACAAAG	2040
CTAGGTGGGA TTATTCGCT ATGAAATGAG AAATTATGGG AGCAAGCTCC TAAATCAACT	2100
GTTTTGATC TACTCTTTA ACTACTTGAT AAAAGTTATA GAAGTAGGCC AAACCTTGAAA	2160
TGATGGTTAC GACTAGGAAT ATTGAAAATT TCCATTGGAC AGGGTTGGTT AAAAGTTGTG	2220
GAAAGGATAT GAGGAGAAAG AAGAGGGCTG CGTTGAGGAC AGGTATCCGT TTTGATTGTA	2280
TTTTCTCAAG TCCTTATTG AGGCCAGGAA GAAAGAGGAG TAGGAGTAGT AAAACTGTAT	2340
GAGAAATAGC TCCTGAAGTA AGGGCGAAGA AAAGAAAAT ACTGATAAAA ACATGAATGA	2400
TCAGTAGTCT AGCTAGTGAT TTCATAAGGC ACCTCCTAAT CCTGGCTTT TTTAGCTCTT	2460
GCAATACGAA GTGAGTCGAC AATATGTATC ATCACTCCGA AAAAGAAAGC TCCCAGTATA	2520
GTTTTAAAAA TATGTTTGT ATTTAGAAGA GAACTGATAA AATTTGGATT TTCACTTGT	2580
AGGGTATCAA TGAGTGGAAAT TATAAAAAT ATCACTGTT CATAAATCGA ACCTGCTTTC	2640
AGACCAGGAT AACGTAACG TTTCTTTCT TTTTCATGA GTTTCCTCCT AATCCTCATC	2700
TTGATTTTC TTAGTTTTG CAATGCGACG GGAGATGAGG AACTGTATGC TCGCTCCGAA	2760
GAAAATAGAA CCGAGAATAC TTGATAACACC ATTTCTTATA GTGAGAAGAG AATGAAAATA	2820
GTCCTGACCT TCATCTATGA GTATCCTGAG AAGAGGAGTT ATAAAAAACAA TCCATAGACC	2880
AAAGAACAAA CCTGCTTCA GACCTGGGTA GTGTAGTTGC TTGCTTTCTT TCTCATTCA	2940
CATATCTGGT TCAATGACTG TGATGCCCTGT TTTTTCTATT TGGTAGGTGA CATAGCCAGA	3000

222

AGCGATGAGG	GCAATCACTA	AAATCAGAGG	AGGATAGATT	AGAGCCACTT	CTTGAGGGTA	3060
TTTATAGGCC	AGAAGGAGTG	GAATAAGATT	TCCGAAAATC	ATCAGATAAA	AGAGGATGAT	3120
AAAGACTTGG	TTCCCAATAC	TATCGGCCCT	ACGCCGTTG	TATTCGTCAA	GGGGACCAGA	3180
AATACCGTAT	GTGCGTTTG	TCAGTTTTTC	AGTGAAGGTT	TCTTTTTCA	TGAGTTTGCT	3240
CCTTTTTAA	AAATCTTCCT	CCCAAAAGAG	ACTGTTGAGG	TCAGTTGGA	GGCTGCGGGC	3300
GAGATTGAGA	CAGAGTTCCA	AGGTGGATT	GTACTTGTG	TTTCAATCA	TATTGATAGT	3360
CTGTCTCGAG	ACACCGATAT	CCTTGGCGAG	TTCGAGCTGG	GAAATACCCA	ATTCCCTTGCG	3420
AAATTCTTTC	ACACGATTCA	TCTGTTCTCC	TTCTGTGATT	ATGTCGTATA	TATTGACTA	3480
TATTATAGTC	TTTTAACAT	AAAGTGTCAA	GTATTTTG	CATATTTTT	GAAGAAATAG	3540
TAGTCTCCTT	GTCCTATTG	TCTGACAAGT	GCAAGCTGGT	CGGATTTGTG	GTAAAATAGA	3600
TAAGATATGA	CAAAAGAATT	TCATCATGTA	ACGGTCTTAC	TCCACGAAAC	GATTGATATG	3660
CTTGACGTTAA	AGCCTGATGG	TATCTACGTT	GATGCGACTT	TGGGCGGGAGC	AGGACATAGC	3720
GAGTATTAT	TAAGTAAATT	AAAGTAAAAAA	GGCCATCTCT	ATGCCTTTGA	CCAGGATCAG	3780
AATGCCATTG	ACAATGCGCA	AAAACGCTTG	GCACCTTACA	TTGAGAAGGG	AATGGTGACC	3840
TTTATCAAGG	ACAACCTCCG	TCATTTACAG	GCATGTTGC	GCGAAGCTGG	TGTTCAAGGAA	3900
ATTGATGGAA	TTTGTATGA	CTTGGGAGTG	TCTAGTCCTC	AATTAGACCA	GCGTGAGCGT	3960
GGTTTTCTT	ATAAAAAGGA	TGCGCCACTG	GACATGCGGA	TGAATCAGGA	TGCTAGCCTG	4020
ACAGCCTATG	AAGTGGTGA	CAATTATGAC	TATCATGACT	TGGTTCGTAT	TTTCTTCAG	4080
TATGGAGAGG	ACAAATTCTC	TAAACAGATT	GCCCGTAAGA	TTGAGCAAGC	GCGTGAAAGTG	4140
AAGCCGATTG	AGACAACGAC	TGAGTTAGCA	GAGATTATCA	AGTTGGTCAA	ACCTGCCAAG	4200
GAACTCAGA	AGAAGGGCA	TCCTGCTAAG	CAGATTTCC	AGGCTATTG	AATTGAAGTC	4260
AATGATGAAC	TGGGAGCGGC	AGATGAGTCC	ATCCAGCAGG	CTATGGATAT	GTTGGCTCTG	4320
GATGGTAGAA	TTTCAGTGAT	TACCTTCAT	TCCTTAGAAG	ACCGCTTGAC	CAAGCAATTG	4380
TTCAAGGAAG	CTTCAACAGT	TGAAGTCCA	AAAGGCTTGC	CTTTCATCCC	AGATGATCTC	4440
AAGCCCAAGA	TGGAATTGGT	GTCCCCTAAG	CCAATCTTG	CAAGTGCAGA	AGAGTTAGAA	4500
GCCAATAACC	GCTCGCACTC	AGCCAAGTTG	CGCGTGGTCA	AAAAATTCA	CAAGTAAGAG	4560
GGAAAAAGAT	GGCAGAAAAAA	ATGGAAAAAA	CAGGTCAAAT	ACTACAGATG	CAACTTAAAC	4620
GGTTTTCGCG	TGTGGAAAAAA	GCTTTTACT	TTTCCATTGC	TGTAACCACT	CTTATTGTAG	4680
CCATTAGTAT	TATTTTATG	CAGACCAAGC	TCTTGCAAGT	GCAGAATGAT	TTGACAAAAA	4740
TCAATGCGCA	GATAGAGGAA	AAGAAGACCG	AATTGGACGA	TGCCAAGCAA	GAGGTCAATG	4800

223

AACTATTACG TGCAGAACGT TTGAAAGAAA TTGCCAATTC ACACGATTTG CAATTAAACA	4860
ATGAAAATAT TAGAATAGCG GAGTAAGATA TGAAGTGGAC AAAAAGAGTA ATCCGTTATG	4920
CGACCAAAAA TCGGAAATCG CCCGCTGAAA ACAGACGCAG AGTGGAAAA AGTCTGAGTT	4980
TATTATCTGT CTTTGTCCCCC GCCATTTTT TAGTCATTT TGCGGTCAATT ATTGGGACAG	5040
GCACTCGCTT TGGAACAGAT TTAGCGAAGG AAGCTAAGAA GGTCATCAA ACCACCCGTA	5100
CAGTTCTGC CAAACGTGGG ACTATTTATG ACCGAAATGG AGTCCCGATT GCTGAGGATG	5160
CAACCTCCTA TAATGTCTAT GCGGTCTATTG ATGAGAACTA TAAAGTCAGCA ACGGGTAAGA	5220
TTCTTACGT AGAAAAAACAA CAATTTAACAA AGGTTGCAGA GGTCTTCAT AAGTATCTGG	5280
ACATGGAAGA ATCCTATGTA AGAGAGCAAC TCTCGCAACC TAATCTCAAG CAAGTTCCCT	5340
TTGGAGCAAA GGGAAATGGG ATTACCTATG CCAATATGAT GTCTATCAAA AAAGAATTGG	5400
AAGCTGCAGA GGTCAAGGGG ATTGATTTTA CAACCAGTCC CAATCGTAGT TACCCAAACG	5460
GACAATTTGC TTCTAGTTT ATCGGTCTAG CTCAGCTCCA TGAAAATGAA GATGGAAGCA	5520
AGAGCTTGCT GGGAACCTCT GGAATGGAGA GTTCTTGAA CAGTATTCTT GCAGGGACAG	5580
ACGGCATTAT TACCTATGAA AAGGATCGTC TGGTAATAT TGTACCCGGA ACAGAACAAAG	5640
TTTCCCAACG AACGATGGAC GGTAAGGATG TTTTACACAC CATTCCACC CCCCTCCAGT	5700
CCTTTATGGA AACCCAGATG GATGCTTTTC AAGAGAAGGT AAAAGGAAAG TACATGACAG	5760
CGACTTTGGT CAGTGCTAAA ACAGGGAAA TTCTGGCAAC AACGCAACGA CCGACCTTTG	5820
ATGCAGATAC AAAAGAAGGC ATTACAGAGG ACTTTGTGTTG GCGTGATATC CTTTACCAAA	5880
GTAACATGAA GCCAGGTTCC ACTATGAAAG TGATGATGTT GGCTGCTGCT ATTGATAATA	5940
ATACCTTCC AGGAGGAGAA GTCTTTAATA GTAGTGAGTT AAAAATTGCA GATGCCACGA	6000
TTCGAGATTG GGACGTTAAT GAAGGATTGA CTGGTGGCAG AACGATGACT TTTTCTCAAG	6060
GTTTTGCACA CTCAGTAAC GTGGGATGA CCCTCCTTGA GCAAAAGATG GGAGATGCTA	6120
CCTGGCTTGA TTATCTTAAT CGTTTTAAAT TTGGAGTTCC GACCCGTTTC GGTTTGACCG	6180
ATGAGTATGC TGGTCAGCTT CCTGGGATA ATATTGTCAT CATTGCGCAA AGCTCATTG	6240
GACAAGGGAT TTCAGTGACCC CAGACGCAAA TGATTCGTGC CTTTACAGCT ATTGCTAATG	6300
ACGGTGTCTAT GCTGGAGCCT AAATTTATTA GTGCCATTAA TGATCCAAAT GATCAAACGT	6360
CTCGGAAATC TCAAAAAGAA ATTGTGGAA ATCCTGTTTC TAAAGATGCA GCTAGTCTAA	6420
CTCGGACTAA CATGGTTTG GTAGGGACGG ATCCGGTTTA TGGAAACCATG TATAACCACA	6480
GCACAGGCAA GCCAACGTAA ACTGTTCCCTG GGCAAAATGT AGCCCTCAAG TCTGGTACCG	6540

224

CTCAGATTGC	TGACGAGAAA	AATGGTGGTT	ATCTAGTCGG	GTTAACCGAC	TATATTTCT	6600
CGGCTGTATC	GATGAGTCGG	GCTGAAAATC	CTGATTTAT	CTTGTATGTG	ACGGTCCAAC	6660
AACCTGAACA	TTATTCAGGT	ATTCAGTTGG	GAGAATTGTC	CAATCCTATC	TTGGAGCGGG	6720
CTTCAGCTAT	GAAAGACTCT	CTCAATCTTC	AAACAACAGC	TAAGGCTTAA	GAGCAAGTAA	6780
GTCAACAAAG	TCCTTATCCT	ATGCCTAGTG	TCAAGGATAT	TTCACCTGGT	GATTTAGCAG	6840
AAGAATTGCG	TCGCAATCTT	GTACAACCCA	TCGTTGTGGG	AACAGGAACG	AAGATTTAAA	6900
ACAGTTCTGC	TGAAGAAGGG	AAGAATCTTG	CCCCGAACCA	GCAAGTCCTT	ATCTTATCTG	6960
ATAAAGCAGA	GGAGGTTCCA	GATATGTATG	GTGGACAAA	GGAGACTGCT	GAGACCCPTG	7020
CTAAGTGGCT	CAATATAGAA	CTTGAATTTC	AAGGTTCGGG	CTCTACTGTG	CAGAAGCAAG	7080
ATGTTCTGTC	TAACACAGCT	ATCAAGGACA	TTAAAAAAAT	TACATTAACT	TTAGGAGACT	7140
AATATGTTA	TTTCACATCG	TGCTGGAATT	GTGACATTTT	TACTAACTTT	AGTAGAAATT	7200
CCGGCCTTTA	TCCAATTAA	TAGAAAGGGC	CAAATTACAG	GCCAGCAGAT	GCATGAGGAT	7260
GTCAACACAGC	ATCAGGCAA	AGCTGGGACT	CCTACAATGG	GAGGTTGGT	TTCTTGATT	7320
ACTTCTGTTT	TGGTGCTTT	CTTTTCGCC	CTATTTAGTA	GCCAAATTCAG	CAATAATGTG	7380
GGAATGATTT	TGTTCATCTT	GGTCTTGTAT	GGCTTGGTCG	GATTTTTAGA	TGACTTTCTC	7440
AAGGTCCTTC	GTAAAATCAA	TGAGGGGCTT	AATCCTAACG	AAAAATTAGC	TCTTCAGCTT	7500
CTAGGTGGAG	TTATCTTCTA	TCTTTCTAT	GAGCGCGGTG	GCGATATCCT	GTCTGTCTT	7560
GGTTATCCAG	TTCATTTGGG	ATTTTCTAT	ATTTTCTTCG	CTCTTTCTG	GCTAGTCGGT	7620
TTTTCAAACG	CACTAAACTT	GACAGACGGT	GTTGACGGTT	TAGCTAGTAT	TTCCGTTGTG	7680
ATTAGTTCTG	CTGCCTATGG	AGTTATTGCC	TATGTGCAAG	GTCAGATGGA	TATTCTTCTA	7740
GTGATTCTTG	CCATGATTGG	TGGTTTGCTC	GGTTTCTTCA	TCTTTAACCA	TAAGCCTGCC	7800
AAGGTCCTTA	TGGGTGATGT	GGGAAGTTTG	GCCCTAGGTG	GGATGCTGGC	AGCTATCTCT	7860
ATGGCTCTCC	ACCAAGAATG	GACTCTCTTG	ATTATCGGAA	TTGTGTATGT	TTTGAAACA	7920
ACTTCTGTTA	TGATGCAAGT	CACTTATTTC	AAACTGACAG	GTGGTAAACG	TATTTCCGT	7980
ATGACGCCCTG	TACATCACCA	TTTGAGCTT	GGGGGATTGT	CTGGTAAAGG	AAATCCTGG	8040
AGCGAGTGGA	AGGTGACTT	CTTCTTTGG	GGAGTGGGAC	TTCTAGCAAG	TCTCCTGACC	8100
CTAGCAATT	TATATTGAT	GTAAGAATGG	CACCCGTATG	TTTCAGGG		8148

(2) INFORMATION FOR SEQ ID NO: 12:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 9909 base pairs
 - (B) TYPE: nucleic acid

225

(C) STRANDEDNESS: double
 (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12:

TAATCCACCC TAAATATCCG TTCTGTAAA TACTTACCG CTTTAAGTT CATAGAATTG	60
AACTTTAAA TGCTTGTCTT CAAGCATCTT TTCCATCAA TTTTTAGGAG TTTGACCAGC	120
TTTAAATAAA AACCTTGCTG GGGTGATTAG TATAGATTTA TCTGCGATTT TATAAGCTTC	180
ATCAATAAAA TAGTGATATA TCGGCTCATC TCTGGCTTCT CCTGTTTCCT GATACGGAGG	240
ATTTCCATTC ACGACATCAA ATTCATTTC ACTTTCCCTCG CTAGATAGGC GCTCAAAACC	300
TATCATTCATA TTCTTTTCC AGTCTTGAT ATGGGTTTA GATTCTTCTA CTTCTTGGAC	360
TTCTAGCTCA TCCGAAACAA AACTCAATTG TTGAGATTGC TTTGTTTAG CTGAATAAGG	420
ACTACTTTT TTCAATCCAT CCATCTGAAA GACATTGTA GAGATAATAG TCGCAATTTC	480
TTTCTTTGC TCTAATGTTG GTGATTTCC AGTCTTAGCT AGATAATAGT CCTCAAAAGT	540
TGCCAAAAGA TTCTCACGCG CCAAAAGGAG AGAATCTCCT TGATACTCCT AACCATACGA	600
AGCATGATAA GCATCTTTA CAAGTTATA AAATGTGACT TCATCTGAAA CCTCACGACT	660
AATCCGTTGC AGTTTTCTAT CAACAAAACC AACTCGCTCA GATAATGGAA TTTCCTCACC	720
AGTTACGGTA TCATATCTCG TTACCATATA AGGTGCTTCA CCACAAGTTA CCTCTAACCA	780
TCGTAAGTCC ACATACTCCT CACACATTAA CGAGCCTAAT TTGATTTCTA CATACTCATT	840
TTGCTTGCG ACCAACACG TTGGTGTAAA CACTTCTGCC CTTATTTTG TCCGATCTTT	900
TTGTTCATAT TTGGATTTTT CAGATCTGGG CTGAATCAAG TTGGCAAAGT TTCCAGTAAC	960
CTTACTTGGA TTGATGCGAT CACTTGGAGC AAATCCCTTT CCTAACAAATT CATAAGAATG	1020
CGTAnGCCAA ACAATTGATT TCTTGTCTG TCGATCTTT AAAAGAATT TTAATAAGTC	1080
AGCCGATTCT TTAGCCAAAC TTCTTCACT AATATCTATT GTCATCAGCA ACCTCTTTA	1140
TATGTAAAGC CCTATTATAT CATATTTAA AGAATGAAAA TTTACTTGAA AAAAGTAATT	1200
CAATAATAT CTCTCGATG ACCAACTTCT AGAGTAGCAA CGACTAATTC ATCATCTACA	1260
ATTTGTACGA TAACTCGATA ATTACCAATT CTATAGCGCC ATTGACCAAC GCGATTACCA	1320
ACCAAAGCCT TTCCGTGTG TCTTGGGTCT TCCAAAACAT TGGTTGTAA ATAGTTGTA	1380
ATTAGCTTCT GCGTATAACG GTCCAATTTC TTCAATTGCT TGATAAAACG TCTTGTGGA	1440
ACTAATTAT ACAAAATTATT CATCCTTCAA GCCTAAATCA TGCATCATT CTTCCAAGT	1500
AATGGGTTCA ACTCCTTTT CCAAGTCTTC TAAATACTCT TGATAGGCTA AATCTGCCAC	1560

226	
ACGAGCATCG TATTCACTTT CTAGGGCTTC AAGAGTTTG GTGCGAATAA GTTCCGAAAG	1620
GGAAACTCCT TCAAACCTAG CCATTGCTTT CATAAATGTT TTATCAGCTT CAGAAACTTT	1680
TAATGTAATA GTAGTCATCT TTTGTGCTCC CTTTTTTAAT GGTAAACACCA TTGTATTACT	1740
TTTTAGGTGT TCAGTCAATA TAAAAAGAAC ACCTTCTCAG CGTTCTTCT ATATCTCTGT	1800
CAATGGTGT GCGGTATCTG GTGAGGTATC ATAAACCTTA AAGTCTACTC CGACTCCCAG	1860
ATCAGCTTGA GCCAGCTGAT TGACCATGGT CATATGAGCC AGTTCCCTGA TATTGTTTC	1920
CTTAGATAAA TGCCCAAGGT AAATCTCTT AGTACGATT CCTAGCGTCC GAATCATAGC	1980
TTCAGCACCG TCCTCGTTAG AAAGGTGACC AAGGTCAGAT AGGATTCCGT GTTGAGTCG	2040
CCAAGCGTAA GAACCTGATC GCAAAATCTC TACATCATGG TTGGCCTCGA TAAGATAACC	2100
ATCCGCATTT TCGACAATGC CCGCCATACG GTCACTGACA TAACCTGTAT CTGTCAGAG	2160
GACAAAATC TTATCATCCT TCATAAAGCG ATAGAACTGC GGTGCGACTG CATCATGGCT	2220
TACACCAAAA CTCTCGATGT CGATATCTCC AAAGGTTTG GTTTTACCCA TTTCAAAAAT	2280
ATGCTTTGC GAAGAATCCA CCTTGCCAAG ATATTTACTA TTTTCCATAG CTTGCCAGGT	2340
CTTTTCATTG GCATAAAGAT CCATACCATA CTTGCGAGCC AAAACGCCA CTCCATGGAT	2400
ATGATCTGAA TGCTCATGGG TAATCAAGAT GGCATCCAGG TCTTCTGGCT TACGGTTAAT	2460
TTCAGCTAGC AGACTGGTAA TTTTCTTGCC AGACAAGCCT GCATCTACTA AAAGCTTCTT	2520
TTTTGAGGTT TCCAGATAAA AAGAATTTC ACTGGAACCC GACGCTAAAA TACTGTATTT	2580
AAAGCCTATT TCACTCATTC TAGTCTCTA CTTCATCCTC CCATACTTCT TCTTTCACTG	2640
CATCCTTATC ATAAGGGAGT ACAATGGTAA AGGTTGAACC CTTGCCGTAT TCACTCTTGG	2700
CCCCAAATAAA GCCCTTATGT TGGTTGATAA TTTCTTTAGC GATAGACAGT CCTAGACCTG	2760
TACCACCTTG TGCACGACTC CTAGCACGAT CCACACGATA GAAACGGTCA AAGATACGTG	2820
GTAAAATCCTG CTTAGGAATC CCCAAACCGT GGTCAGAAAT GGATAAAATC ATCTGGCTT	2880
CAGTTGTCTT CATTCTGACA GTGATTTAC CCCCCATCTGG CGAATACTTA ATAGCATTAT	2940
TTAAAATATT GTCGACAACC TGGTGTATCT TATCTGTATC AATTTCATC CAGATAGAAT	3000
TGATGGGATA ATCTCTCACC AACTCATATT TTTCTCCTT TTCCTGTCCT TTCATCTTGT	3060
CAAAACGATT GAGGATAAAAG GTAATAAAAG CAGTGAAGTT AATCAGTTCC ACATCTAGGT	3120
GAATGGTAGC ATTATCAATA CGTAAAGAT GGAGGAGATC CGTCACCATG CGCATCATAAC	3180
GGTTGGTCTC ATCAAGAGAA ACCTTGATAA AGTCTGGTGC TACAGTTCA CACAAAGCCC	3240
CCTCATCCAA GGCTTCAGA TAGGATTTA CGCTAGTCAG AGGAGTCCGT AACTCATGGC	3300
TAACATTGGA AACAAAGAGT CTTCGTTCGC GTTCTTCCTT CTCCGTCTCC GTCGTATCAT	3360

227

GCAAAACAGC CACCAAACCT GAAATAAAGC CAGACTCTCG ACGTATCAAG GCAAAGCGAA	3420
CTCGAAGGTT CAAATATTCG CCATTGATAT CTTGGGAATC TAGCAACAAT TCTGGACTTT	3480
GGGTAATCAA ATCACGCAAT TCATAGTTT CTTCTATCTT GAGCAATTCC AAAATGCTTC	3540
TATTCAGAAC ATCTTCCTTA ACCAACCCCA GTTGCTTCTT GGCTGTATCG TTAATCATGA	3600
TAATCTGACC CCGACGGTTA GTCCAAGAA CCCCATCTGT CATATAAAAC AGAATACTAT	3660
TTAGCCTCTT ACTCTCTTGT TCTAGATTTT CCTGAGTGTAG ACGAATAACC TCCGACAAGT	3720
CATTCAAATT ATTGGTAATA TTGGTGATTT CAGACCCACC TTGCATATCA AGAACCTTGG	3780
AATAATCTCC TGCAATCAA TCTTTAACCT TTGATTGAC TTGCTTCAAC TGAATATTAT	3840
CACGTCTATT TTCCAGTAAT AAGAGGGTCA CAACAAGGAT GAAACCTAAC AAAATCAGGA	3900
TAAAGATAAA ATCTCTGGTA AAAATGGTTT GTTTCAGTAA ATCAAGCATT ATTTCTCATG	3960
TAATACCCCTA CACCACGGCG CGTCAAGATA TACTCTGGTC GGCTGGCGT ATCTTCATC	4020
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CCCCAGACAG TCTCAAGCAA GTGTCGCGC GTGATGACTT GACCTGTATG CGATGCTAAA	4140
TGATACAAAA GCTCAAATTC ACGATGGTTT AAGTCTAGTT CTTCGCCATA TTTTTAGCC	4200
ACGTAGGCCTGCTGGAAATTTCC CCAATTGGAA TAGGTTGAGG TTTACTATCT	4260
GCTTCCTGAC CATCTACTGG CATAAGTTGA GAACGACGCA GAAGAGCTTT AACACGCGCC	4320
TGCAACTCAC GATTGGAGAA GGGTTTGTT ACATAGTCAT CTGCCCAAG TTCCAACCG	4380
ATAACCTTAT CAAATTCACT ATCTTTGGCT GAAAGCATAA GAATGGGCAC ACTGCTTGTC	4440
TTACGAATGG TCTTAGCAAC TTCTAAACCA TCAATTCTG GAAGCATCAA ATCCAGAATA	4500
ATAATATCTG GTTGCTCTGC TTCAAATTGC TCTAGCGCTT CACGACCATT AAAAGCAGTT	4560
ACAACCTCGT AACCTCCCTT GGTCATATTA AACTTGATAA TATCCGAGAT TGGTTCTCA	4620
TCATCTACAA TTAGTATTTT TTTCATATGT TCACCTTTT CTCTACTATT ATACCAAAA	4680
AATAGTCAGA AGACACAATA GCTAGCTTG GCTACTGTCT AAGTTGGCTT GTGCATAAAC	4740
CTGCCAGATT TTTTGGGG GTTGGCAAG TGGGTAATTG TTGAATTCTT CTGGTAAAG	4800
CCAGCGAACT TCCCTATCTG AAAAATCATG GAAGTCACTC ACCTGACCTG CTACAATCTG	4860
TACATGCCAT TTTCGATGAC TAAAAACATG CTGGACTGTA TCAAAACAAA CATCAAGCCA	4920
ATCAACATCT AGGTCAAGT CCTGCTGGAA ACTCTCTCT GGACTGGAC CAAAGTTCAC	4980
ACTTTCTTCC GCAACCTGAT GAAAGAGGTC AACTGCTCT TCTTGCGAAA AGTTATCAAC	5040
TTCTATAAAG GGGAAATGCC AAAAACCTGC CAAGAGCTTT TCGCTTCAT TTTTTCAAG	5100

228						
TAAAAATTGT	CCTTGAGAAT	TTTCACAAC	TAAGGTTTA	AGATAAATAG	GAACCGGCTT	5160
TTTCTTAGGA	GATTTAATTG	GATAACGGTC	CATGGTCCA	TTCTGATATG	CCGCACTAA	5220
GTCCCTGACT	GGGCTTCCTT	CAGGTCTGGG	ATTACAGGA	GAETCAATAT	CAGACCCCTAA	5280
GTCCATCAAG	GCTTGATTAA	AATCACCCGG	ACGATCCGGA	TTAATCAAGA	TCTCCATCAT	5340
TGCCCTGAAAA	ATTTTCGAT	TACTTGAAT	CCCAATATCG	TGGTTGACTT	CAAACAGACG	5400
CGCCAAGACC	CGCATGACAT	TACCATCTAC	AGCTGGCTCA	GGCAAGTTAA	AAGCAAACT	5460
GGAAATGGCT	CCTGCTGTGT	AAGGTCCAAT	CCCTTCAAG	CTGGAAATTC	CTTCATAGGT	5520
ATTTGGAAT	TGGCCACCAA	AGTCAGTCAT	AATCTGCTGG	GCTGCAGCCT	GCATATTGCG	5580
AACTCGAGAA	TAATAGCCCA	AGCCCTCCC	AGCTTTCAGT	AAACTCTCCT	CAGGCGCAGT	5640
TGCCAGACTT	TCGACAGTTG	AAAACCAGTC	AAAAATCTT	TCGTAGTAAG	GGATAACTGT	5700
ATCCACCCCTG	GTCTGCTGAA	GCATGATTTC	AGATAACCCAG	ATGTGATAAG	GATTTTTACT	5760
TCTCCTCCAA	GGCAAATCTC	TTTGTTTTC	ATCATACCAA	GCGAGAAGTT	TCTCACGGAA	5820
AGAAATGACT	TTCTCCTCCG	GCCACATGAC	GATAACGTAT	TCTTCAAAAT	CTAACATATC	5880
TCTAGTATAA	CACAGAAGGT	TTCACCTGTC	TTTGTATCTG	ATTTATAATA	TTTTCAATAG	5940
ATAGTATATA	ACTTTCTAT	CTACTTATAC	TCAATGAAAA	TCAAAGAGCA	AACTAGGAAG	6000
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ATCATATACt	ACGGCAAGGT	GAAGCTGACG	TAGTTGAAG	AGATTTTCGA	AGAGTATAAA	6120
TCTTATTGAT	GAACTGCTTG	CAGTCTGAGA	AAAAATGAGC	TTGGATATTA	TTTCCAAACT	6180
CACTTAAAGT	CAATTCAAT	CCACTAGAAC	AAGCCTAGTA	CAGTCCATC	GCTTCAACA	6240
TCCATGTTGA	GAGCTGCTGG	ACGTTTTGGA	AGACCTGGCA	TGGTCATAAC	ATCACCAGTT	6300
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CAGATTGGCA	ATTTGTCCC	ACCGTTTGAG	ACGATTGAG	CAATTTGTGT	TTGAGCTTTC	6480
TTCTCAAAGT	TCACTTTGCT	ACCACGATAG	ATTCAGTGA	CAATTTTTTC	AATCTTTCT	6540
TGGACAGAAA	GGTCATTATC	ATACAAACGT	TTATAGTTAG	CTGGATTTTC	AGCAATTGTC	6600
TTAACAACTG	TTTCGGCAAG	TGCTACTCCA	CCTCTGCTC	CATCAGCCCA	GACACTAGCC	6660
AATTCAACTG	GTACATCGAT	TGAGGCACAG	AGTTCTTTA	AGGCTGCAAT	TTCAGCTTCT	6720
GTATCAGATA	CAAATTGTT	AATAGCTACA	ACTGCTGGAA	TACCGAACTT	ACGGATATT	6780
TCAACGTGGC	GTTCAAGTT	AGCAAAACCT	GCACGAACTG	CCTCTACATT	TTCTTCAGTC	6840
AGAGCGCTT	TAGCCACACC	ACCATTCAATC	TTAAGGGCAC	GAAGGGTTGC	GACAATAACA	6900

229

ACTGCATCTG GAGATGTTGG CAAGTTGGT GTCTTGATAT CAAGGAATTT CTCAGCACCA	6960
AGGTCCGCAC CAAAACACAGC TTCACTAACCA GTGTAATCAG CCAAGTGAAG GGCTGTTGTC	7020
GTCGCCAAAA CAGAGTTACA GCCATGAGCG ATATTGGCAA ATGGACCACC GTGTACAAAG	7080
GCAGGGTGTAC CGTAAATTGT CTGAACCAAG TTTGGCTTAA TAGCATCCTT CAAAATCAA	7140
GCCAAGGCAC CCTCAACCTG CAAATCACCT ACAGAAACAG GCGTACGGTC ATAGCGATAA	7200
CCAATAACGA TATTCGCCAA ACGACGTTTC AAGTCCTCGA TGTCGGTTGC CAAGCAAAGA	7260
ATTGCCATGA TTCTGAAGC AACTGTAATA TCAAAACCAT CCTCACGTGG AATACCGTTT	7320
AGAGGACAC CAAGACCAAC AGTCACATGG CGGAGCGTAC GGTGTTCAA GTCCACAAACG	7380
CGTTTCCAGA GGATACGACG TTGATCAATT CCCAGCTCAT TCCCTGGTG CAAGTGGTTG	7440
TCAATCAAGG CAGAAAGGGC ATTGGTGGCA GTTGTAAATAG CATGCATATC TCCAGTAAAG	7500
TGGAGGTTGA TGTCTTCCAT TGGCAGAACT TGTGCATACC CACCACCAGC AGCACCACCC	7560
TTGATCCCCA TGACTGGACC AAGAGACGGT TCGCGGATAG CAATCATGGT TTTCTTGCCA	7620
ATCTTGTCA AGGCATCCGC AAGACCAATG GTAAGCGTCG ACTTTCTTC ACCTGCAGGT	7680
GTTGGGTTGA TGGCAGTAAC CAAGATCAAT TTACCGACTG GATTGCTCTC AACTGCACGA	7740
ATTTTATCAA AGCTGAGTTT AGCCTGTAC TTTCCGTACA ACTCCAAATC GTCATAAGAA	7800
ATACCAAGTT TCTCTACAAAC ATCAACAATT GGCTTCAACT CAATACTCTG TCGGATTTCA	7860
ATATCTCTT TCATTCAAAA TTCCCTCAAC CTCTTATATG ATAATTCTATT ATATCACAAA	7920
ACAAGAGTTT TAACATCCTA AAACTCTCTA AACGTTCGTA AATATCTCTG TTTTTAAGAC	7980
TTTTAGAGTC CTTTCTTAAA TTTTATATGG CTTTATAGTT TGAAACTATA ATAAATCTTC	8040
GTTTTTACCA AAAATTATC ACTTTCATTT TACTTACCGC TTATTTTGT GTACAATAGT	8100
GCTATGAAAA TTTTAGTTAC ATCGGGCGGT ACCAGTGAAG CTATCGATAG CGTCGCTCT	8160
ATCACTAACCC ATTCTACAGG TCACTGGGG AAAATTATCA CAGAGACTTT GCTTTCTGCA	8220
GGGTATGAAG TTTGTTTAAT TACGACAAAA CGAGCTCTGA AGCCAGAGCC TCATCCTAAC	8280
CTAAGTATTC GAGAAATTAC CAATACCAAG GACCTCTAA TAGAAATGCA AGAACGTGTT	8340
CAGGATTATC AGGTCTTGAT CCACTCAATG GCTGTTCTG ACTACACTCC TGTTTATATG	8400
ACAGGGCTTG AGGAAGTTCA GGCTAGCTCC AATCTAAAAG AATTTTTAAG CAAGCAAAAT	8460
CATCAGGCCA AGATTCTTC AACTGATGAG GTTCAGGTTT TGTTCCCTAA AAAGACACCC	8520
AAAATCATAT CCCTAGTCAA GGAATGGAAT CCTACTATTC ATCTGATTGG TTTCAAAC	8580
CTGGTTGATG TTACCGAAGA TCATCTGGTT GACATTGCAC GAAAAAGTCT TATCAAGAAT	8640

230	
CAAGCAGATT TAATCATCGC GAATGACCTG ACTCAAATTT CAGCAGATCA GCACCGAGCT	8700
ATATTTGTTG AGAAAATCA GCTTCAAACA GTCCAGACTA AAGAAGAAAT TGCAGAACTC	8760
CTCCTGAAA AAATCAAGC CTATCATTCT TAGAAAGGAA AACTATGGCA AACATTCTCT	8820
TGGCTGTAAC GGGTCAATC GCCTCTTATA AGTCGGCAGA TTTAGTCAGT TCTCTAAAAA	8880
AACAAGGCCA TCAAGTCACT GTCTTAATGA CTCAGGCTGC TACAGAGTTT ATCCAACCTT	8940
TGACACTACA GGTACTCTCA CAGAATCCTG TCCACTTGGA TGTATGAGA GAACCTATC	9000
CTGATCAGGT CAATCATATC GAACTTGAA AAAAGCAGA TTTATTTATC GTGGTACCTG	9060
CAACTGCTAA CACTATTGCA AAACTAGCTC ACGGATTTGC GGACAACATG GTAACCAGTA	9120
CAGCTCTAGC CCTACCAAGT CATATTCCA AACTAATAGC TCCTGCTATG AATACAAAAA	9180
TGTATGACCA TCCAGTAACT CAGAATAATC TGAAAACATT AGAAAACATCG GCTATCAGCT	9240
GATTGCTCCT AAGGAATCCC TACTAGCTTG TGAGGACAC GGACGAGGAG CTTTAGCTGA	9300
CCTCACAAATT ATTTAGAAA GAATAAAAGGA AACTATCGAT GAAAAAACCC TCTAATATTG	9360
CACCCATTGC TATCTTTTT GCTACCATGC TCGTGATACA CTTTCTGAGC TCACCTATCT	9420
TTAACCTTTT TCCATTTCCA ATCAAACCGA CCATTGTTCA TATTCTGTC ATTATTGCCA	9480
GCATTATTAA TGGTCCACGA GTGGGGTTA CACTTGGATT TTTGATGGGA TTACTTAGCT	9540
TGACGGTTAA CACGATTACG ATTCTACCGA CAAGCTACCT CTTCTCTCCC TTCTGACCAA	9600
ACGGAAACAT CTACTCAGCT ATCATTGCCA TCGTCCCACG TATTTTGATT GGTTTAAC	9660
CTTACTTAGT CTATAAACTG ATGAAAACA AGACTGGTCT GATTTTAGCT GGAGCCCTG	9720
GTTCCTTGAC AAATACTATC TTTGTCCCTG GAGGAATCTT CTTCTATTT GGAAATGTT	9780
ATAATGGAAA TATCCAACCT CTTCTGGCAA CCGTTATCTC AACAAATTCA ATTGCTGAAT	9840
TGGTCATTTC TGCAATTCTA ACCCTAGCCA TTGTTCCACG ACTACAAACC TTGAAAAAT	9900
AAAAACAGG	9909

(2) INFORMATION FOR SEQ ID NO: 13:

- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 1126 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13:

TAATTTTCAT ATAATAGTAA AATAGAATGT GTGATTCAAT AATCACCTCA AATAGAAAGG	60
AAATTCTATG TCAAATCTAT CTGTTAATGC AATTCGTTTT CTAGGTATTG ACGCCATTAA	120

231

TAAAGCCAAC	TCAGGTCATC	CAGGTGTGGT	TATGGGAGCG	GCTCCGATGG	CTTACAGCCT	180
CTTTACAAAA	CAACTTCATA	TCAATCCAGC	TCAACCAAAC	TGGATTAACC	GCGACCGCTT	240
TATTCTTC	GCAGGTCATG	GTTCAATGCT	CCTTTATGCT	CTTCTTCACC	TTTCTGGTT	300
TGAAGATGTC	AGCATGGATG	AGATTAAGAG	TTTCCGTC	TGGGGTTCAA	AAACACCAGG	360
TCACCCAGAA	TTTGGTCATA	CGCAGGGAT	TGATGCTACG	ACAGGTCC	TAGGGCAAGG	420
GATTTCACT	GCTACTGGTT	TTGCCCAAGC	AGAACGTTTC	TTGCCAGCCA	AATAAACCG	480
TGAAGGTTAC	AATATCTTG	ACCACTATAC	TTACGTTATC	TGTGGAGACG	GAGACTTGAT	540
GGAAGGTGTC	TCAAGCGAGG	CAGCTTCATA	CGCAGGCTT	CAAAA	ACTTG ATAAGTTGGT	600
TGTTCTTAT	GATTCAAATG	ATATCAACTT	GGATGGTGAG	ACAAAGGATT	CCTTACAGA	660
AAGTGGT	CGT GACCGTTACA	ATGCCTACGG	TTGGCATACT	GCCTTGGTTG	AAAATGGAAC	720
AGACTTGGAA	GCCATCCATG	CTGCTATCGA	AAACGAAAA	GCTTCAGGCA	AGCCATCTT	780
GATTGAAGTG	AAGACGGTTA	TTGGATACGG	TTCTCCAAAC	AAACAAGGAA	CTAATGCTGT	840
ACACGGGCC	CCTCTGGAG	CAGATGAAAC	TGCATCACT	CGTCAAGCCC	TCGGTTGGGA	900
CTACGAACCA	TTTGAATTC	CAGAACAACT	ATATGCTGAT	TTCAAAGAAC	ATGTTGCAGA	960
CCGTGGCGCA	TCAGCTTATC	AAGCTTGGAC	TAAATTAGTT	GCAGATTATA	AAGAAGCTCA	1020
TCCAGAACTG	GCTGCAGAAG	TAGAAGCCAT	CATCGACGGA	CGTGATCCAG	TCGAAGTGAC	1080
TCCAGCAGAC	TTCCAGCTT	TAGAAAATGG	TTTTCTCA	CCAACT		1126

(2) INFORMATION FOR SEQ ID NO: 14:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2520 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14:

CCGGCAACAA	AAAAGAAAAA	ATCAACAGTT	AAAAAAAATC	TAGTCATCGT	GGAGTCGCCT	60
GCTAAGCCAA	GACGATTGAA	AAATATCTAG	GCAGAAACTA	CAAGGTTTA	GCCAGTGTG	120
GGCATATCCG	TGATTGAG	AAATCCAGTA	TGTCCGTCGA	TATTGAAAAT	AATTATGAAC	180
CGCAATATAT	TAATATCCGA	GGAAAAGGCC	CTCTTATCAA	TGACTTGAAA	AAAGAAGCTA	240
AAAAAGCTAA	TAAAGTTTT	CTCCGAGTG	ACCCGGACCG	TGAAGGAGAA	GCGATTCTT	300
GGCATTTGGC	CCATATTCTC	AACTGGATG	AAAATGATGC	CAACCGTGTG	GTCTCAATG	360

232	
AAATCACCAA GGATGCAGTC AAAATGCTT TAAAGAACCC TCGTAAGATC GATATGGACT	420
TGGTCGATGC CCAACAAGCT CGTCGGATCT TGGATCGCTT GGTAGGGTAT TCGATTTCCG	480
CTATTTGTG GAAGAAGGTC AAGAAGGGCT TGTCAAGCAGG TCGCGTTCAG TCCATTGCC	540
TTAAACTCAT CATTGACCGT GAAAATGAAA TCAATGCCTT CCAGCCAGAA GAATACTGGA	600
CAGTTGATGC TGTCTTTAAA AAGGGAACCA AACAAATTCA TGCTTCCTTC TATGGAGTAG	660
ATGGTAAAAA GATGAAACTG ACCAGCAATA ACGAAGTCAA GGAAGTCTTG TCTCGTCTGA	720
CGAGTAAAGA CTTTCAGTA GATCAGGTGG ATAAGAAAGA GCGCAAGCGC AATGCTCCTT	780
TACCCATAC CACTTCATCT ATGCAGATGG ATGCTGCCAA TAAAATCAAT TTCCGTACTC	840
GAAAAACCAT GATGGTTGCC CAACAGCTCT ATGAAGGAAT TAATATCGGT TCTGGTGTTC	900
AAGGTTTGAT TACCTATATG CGTACCGATT CGACTCGTAT CAGTCCTGTA GCGCAAAATG	960
AGGCGGCAAG CTTCATTACG GATCGTTTG GTAGCAAGTA TTCTAACGCAC GGTAGCAAGG	1020
TCAAAAACGC ATCAGGTGCT CAGGATGCCC ATGAGGCTAT TCGTCCGTCA AGTGTCTTTA	1080
ATACACCCAGA AAGCATCGCT AAGTATCTGG ACAAGGATCA GCTTAAGCTA TATACCCTTA	1140
TCTGGAATCG TTTTGTGGCT AGCCAGATGA CAGCGGCCGT TTTTGATACC ATGGCTGTTA	1200
AATTGTCCTCA AAAAGGGGTT CAATTGCTG CCAATGGTAG TCAGGTTAAG TTTGATGGTT	1260
ATCTTGCCAT TTATAATGAT TCTGACAAGA ATAAGATGTT ACCGGACATG GTTGTGGAG	1320
ATGTGGTCAA ACAGGTCAAT AGCAAACCG AGCAACATT CACCCAACCG CCTGCCCGTT	1380
ATTCTGAAGC AACACTGATT AAAACCTTAG AGGAAAATGG GTTGGACGT CCATCAACCT	1440
ACGCCCAAC CATTGAAACC ATTCAGAAC GTTATTATGT TCGCCTGGCA GCCAACCGTT	1500
TTGAACCGAC AGAGTTGGGA GAAATTGTCA ATAAGCTCAT CGTGAATAT TTCCCGAGATA	1560
TCGTAAACGT GACCTTCACA GCTGAAATGG AAGGTAAACT GGATGATGTC GAAGTTGGAA	1620
AAGAGCAGTG GCGACGGGTC ATTGATGCCT TTTACAAACC ATTCTCTAAA GAAGTTGCCA	1680
AGGCTGAAGA AGAAATGGAA AAAATCCAGA TTAAGGATGA ACCAGCTGGA TTTGACTGTG	1740
AAAGTGTGTGG CACTCCAATG GTCAATTAAAC TTGGTCGTTT TGTTAAATTAC TACGCTTGT	1800
GCAATTTCCTT AGATTGCCGT CATAACCAAG CAATCGTAA AGAGATTGGT GTTGGAGTGT	1860
CAAGCTGTCA TCAGGGACAA ATTATTGAGC GAAAACCAA GCGTAATCGC CTATTCTATG	1920
GTTGCAATCG CTATCCAGAA TGTAATTAA CCTCTGGGA CAAGCCTGTT GGTCGTGACT	1980
GTCCAAAATG TGGCAACTTC CTCATGGAGA AAAAGTCCG TGTTGGTGGC AAGCAGGTTG	2040
TTTGTAGCAA AGGCAGTAC GAGGAAGAAA AGATGGCTCT TTGTCAACTG TAGTGGTTG	2100
AAGTCAGCTA AGCTCGAGAA AGGACAAATT TTGTCCCTTC TTTTTGATA TTCAGAGCGA	2160

233

TAAAAATCCG TTTTTGAAG TTTTCAAAGT TCCGAAAACC AAAGGCATTG CGCTTGATAA	2220
GTGATGAG ATTATTGGTC GCTTCCAATT TGGCGTTAGA ATAGTGTAGT TGAAGGGCGT	2280
TGACGATTTT CTCTTTGTCC TTTAGAAAGG TTTTAAAGAC AGTCTGAAA AGAGGATGAA	2340
CCTGCTTTAG ATTGTCCTCA ATGAGTCCGA AAAATTCTC CGGTTCCCTTA TTCTGAAAGT	2400
GAAACAGCAA GAGTTGATAG AGCTGATAGT GATGTTCAA GTCTTGTGAA TAGCTCAAAA	2460
GCTTGTAAAT AATCTCTTAA TTGGTTAAAT GCATACGAAA AGTAGGGCGA TAAAAATGTT	2520

(2) INFORMATION FOR SEQ ID NO: 15:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 10993 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15:

TTTTCTCGAT AATAACTTCC ACCTTATTAT TTGGGATACC CTCTCTTCT TCACCACAC	60
GTTCATAGTA GTCATCGCGA TAGAGAAAAG CTACGATATC AGCGTCCTGC TCAATAGACC	120
CAGATTCACG AATATCAGAC AAGACCGGTC TCTTGTCCCTG ACGTTGTTCT ACACCACGAG	180
AAAGCTGACT CAGAGCGATT ACTGGAACCT TCAATTCTT GGCTAGTATT TTCAACTGAC	240
GAGAAATTTC AGAAACTCT TGTGACGAT TTTCTCGACC AGTCCCGTG ATAAGTTGCA	300
AATAGTCTAT CAAAATCAA CCAAGATTC CAGTTCTTG AGCCAATTAA CGAGAACGAG	360
AACGAATCTC TGTAAATCGA ATACCTGGCG TATCATCGAT ATAGATACTG GCGTTAGCTA	420
GATTACCTG AGCAATAGTA TATTTTGCC ACTCCTCATC TGTCAATTGC CCTGTACGGA	480
TAGAATGTGA CTCCACTAAG CCTCTCTGCAG CTAACATACG ATCTACCAAG CTTTCCGCAC	540
CCATTTCGAG TGAAAAAATA GCAACCGTT TGTCACATT AGTCCCAATG TTCTGAGCGA	600
TATTCAGGC AAATGCTGTC TTACCAACTG CTGGACGAGC TGCTAAGATA ATCAACTCCT	660
CCTCATGAAG TCCCTGTGTC ATATGATCCA AATCACGATA ACCTGTCGCA ATACCTGTAA	720
TATCGGTCGT TTGTTGCGAG CGAGCTTCCA GATTTCCAAA GTTGAGATTC AACACATCTC	780
GAATGTTCTT AAACCCGCTT CGATTTGCAT TTTCACTGAC ATCAATCAAC CCTTTTCTG	840
CCTGAGCAAT AATTTCATCA GCTGGTTGTG ACGCTTCGTA AGCTTGGTG ACAGACTCTG	900
TCAACTTGGC AATTAACGA CGTAGCATTG CTTTTCTGC AACAAATCTTA GCATAATACT	960
CCGCATTAGC AGAAGTTGGC ACAGAATTAA CAATCTCAAC CAAGTAAGAC AAGCCACCAA	1020

234	
TATTCTGTAA ATCACCTGAA TTATCAAGGA TAGTACGAAC CGTTGTTGCA TCTATGGCAT	1080
CACCAACGATC GGATAAAATCG ACCATGGCTT GGAAAATCAA ACGATGGGCA TACTTAAAAA	1140
AGTCCCGAGA CTCAATGTAT TCTCGCACAA AAACAAGTTT ACTCTCATCA ATAAAGATAG	1200
CCCCTAAAC GGATTGCTCA GCTAAGATAT CTTGAGGTTG TACTCGTAAC TCTTCTACTT	1260
CTGCCATCACTG ACTTCCCTTC CTTTACAAT CTTGTCAAGA AGGTGTAAAC TTATCCTTCT	1320
TTCACACGAA GATTGATTAC ACTTGTGATA TCTTGATAGA TTTTCAGTGG CACATCAATC	1380
AAACCAACCG CTCGAATCGG AGCTTGTACT TGAATATGAC GTTTATCAAT CTTAATTCCA	1440
AATTGCTTTT GCAATTCTTC TGCAATCTTC TTATGGTAA TAGAACAAA GGTACGACCA	1500
TCTGGACCAA CTTTTCAAC AAATTCTACA ACAGTTCTT CTGCTTCAAG TTGTGCTTTA	1560
ATTGCTTTTC CTTCTGCAAT CATCTCAGGG TGAGCTTTT CTTCCGATTT TTGTTTACCA	1620
CGAAGTTTCAC CTACAGCTTG AGCAGTCGCT TCTTGGCTA GATTCTTTT GATAAGAAAG	1680
TTTTGCCAT ACCCTGTTGG TACTTCCTTA ATTTGCCCTT TTTTACCTTT TCCTTTAACAA	1740
TCTGCTAAA AGATTACTTT CATTCTCTT TCTCCTTTTC CTTCATTTCA TTTAATACAA	1800
TTTCTGTCAG TTTTCACCT GCTTCTGACA AGGTTACATC TTTAATTGAA GCTGCTGCCA	1860
AATTAAAGTG GCCTCCACCG CCTAACTCTT CCATAATCGG TTGTACATTC AGTTTACTAC	1920
GACTTCGAGC TGAGATAGAG ATAATCCCTT GTGTATTCTT CGCAAGAACAA AAACCTCGCTT	1980
CAATACCTGA CATGGCTAAC ATGGCATCTG CTGCCCTACT AATAACAACAT GTATCATAGC	2040
ATTTCATGTC CTTAGCCTCT GCTATTAGTA CATCTGAACC TAATTTACGC CCCTGTAAAAA	2100
TAAGTTCAATT GACCTCACGA TATTCTCAA AATCTGTCGC AGCGATTTCG TGGATAGCAA	2160
TACTATCACT TCCGCGCGTT CTGAGATAGC TAGCAACATC AAATGTCCGA CTAGTTACTC	2220
GCGAGGTGAA ATTTTAGTA TCCAACATCA TACCAGCCAT CAAGACACTT GCTTGCATAC	2280
GACTCAAACG ATTTCTTAA GAATTCTGGA ACTGAATCAA TTCCGTTACC AACTCACTGG	2340
CACTACTTGC ACCACTTCG ATATAAGTAA TAACCGCATT ATCTGGAAAA TCCTGATCCC	2400
TTCTATGGTG GTCAATAACA ATGGTTGGG TAAATAAATC ATAAAATTCT TTTGATAATG	2460
TTAAGGCTGT CTTGAATGG TCTACAAGAA TCAACAAAGA ACGATTGGTC ACCATCCCCA	2520
TTGCATCCTT AACAGACAAC AACCTCGTAA CTCCCTCTT TTCTATGAAT GAAACAGCTC	2580
GTTCAATATC TGGAGACATT TGTTCTTCAT CATAAAGAGC ATAGCTATTT TCAATCACAT	2640
TGCTGGCGAA CAACTGCATA CCTACAGCAG AGCCCAAAGC ATCCATGTCT AAATTTTGT	2700
GACCGACTAC AAAACCTGA TCTACACTCC GAATCTTATC TGAAATAGCT GTCATCATAG	2760
CGCGCGTACG AGTCCGTGTA CGCTTGATTG AAGCAGCAGA CCCACCACCA AAATAAACTG	2820

235

GATTTTCGTTTTCGTTTTCCTTAACAA	2880
CCACCTGGTCGCCACCACGTACTTCAGCCA	
AGTTCAAATTGAGCAAAGCACTTTCCCTATCTCATCATG	2940
ATTTCCATCGCCATAAGAAA	
ATCCCATACTTAAGGTCAAGGGCAACTGTC	3000
TCTGTTTCGACTCTCTCTGAAACCATCAA	
TAACAGAAAAA	
TTTATCATTCATCAAGGCCCTCAAGCACCGGTG	3060
TAGTCAGTAATAAGATAAA	
ATCGATCCATACTTACCCGA	3120
CGAGAAAACA	
TCATGTGTTTTTCTCTGAAAC	
TCTGATATAAA	
AATTAGCTACAAA	3180
ACTTACCCGATTTGACTAA	
TATCTGACTCAGAAGTTTCATCCCTAA	
TCCTCCAAAT	
CATCATAATTATCCACAGAG	3240
ACAATCCCAA	
TCACTGGTCTACTTGTTACC	
AATTCACTG	
TTATGGCTTGTTCCCTGGAT	3300
ACATCTACAA	
AATACAAAAC	
ACCGGAAGAA	
GCATCCATAT	
GAACAGCATAACGCTTCTCA	3360
CCAAGCTTGGCATAAGTAGA	
CGGATTTCCACTGAAGCCT	
TGATAATCGTTGAAACAGCTCTAAATCAA	3420
AATCACCAC	
TTCCCTGGTC	
AAAATCAATT	
CAGCATAGGG	3480
ATTAACCAC	
TCAACCTCTC	
CAGAAGATAA	
ATTCAATTTC	
ATAACACCTA	
CAGGCATCTGTTCCAAATAGA	3540
GCTGTCAAC	
TTTCTTCCGC	
TTGGTGGTTTACATACTGTA	
TCTGTTCTAC	3600
ATCACTCCTT	
GTATAATGCA	
CTCTCAGTTTT	
CTTAAATAAA	
AAAACATAGC	
CTCCTACAA	3660
AAGAACACAA	
ATTAAACCG	
TCAACAGATT	
ATTATTAACA	
AAAATAATGA	
AAGTGGATAA	3720
GACTCCAAAC	
GCAATCAATC	
CTACTAGAAT	
AGGAAAATT	
GGACTTACAT	
AAAATTTTT	3780
CATTCAAAAC	
CTCTTGGCAC	
CCATTATACC	
ATAATACCCC	
TCAAAAGCG	
ACTTTTTAAGA	3840
AGTGTAAATCA	
GTAATTCTAT	
CAATTATAAG	
AAAAGGTAG	
TTTACAATT	
AGTAAACCTA	3900
CCTTACACAA	
TATTGAAATT	
AAGATTCTTT	
AACCTCTAAC	
AAACCAATT	
CGCCATCCTC	3960
ACGACGATAA	
ATCACATTGG	
TTGTCTGATC	
TTCAACATCC	
ACATAGATAA	
AGAAATCATG	4020
CCCCAAATAAA	
TCCATTGTA	
GAATTGCTTC	
TTCCAATTC	
ATTGGTTTTA	
AATCAATTG	4080
TTTGAAACGA	
ACAACTTAG	
ACTGGACAA	
ATTGAAATCT	
TCCACCAAAG	
CATCTGAA	4140
TAATTGACCA	
GTTGCTACCT	
TATTTTTATT	
TTTACGCTCG	
ATTTTTGTTT	
TATTTTACG	4200
AATCTGACGT	
TCAATTCTAT	
CAGTTACAAG	
GTCAATTGAA	
CCATACATAT	
CTTGAGATAC	4260
ATCTCTGCG	
CGGAGAGTAA	
TAGATCCAAG	
CGGAATCGTT	
ACTTCCACTT	
TAGCCGTTTT	4320
TTCACGATAA	
ACTTTTAAGT	
TAATTCGGGC	
ATCCAACCT	
TGTTCTGGTT	
GGAAGTACTT	4380
TTCGATCTTT	
TCGAGTTAG	
AAACTACACATA	
ATCACGAATT	
GCTTCTGTTA	
CTTCTAGGTT	4440
TTCACCAACGG	
ATACTATATT	
TAATCATATG	
AGTACCTTCT	
TTCTAAACAT	
TTTTGTTTTT	4500
ATGATTTAT	
TATAACGCTT	
TCATTCTATT	
TTTGCAAATT	
TTTCCTCAT	
CTTACAAGGG	4560
AAAATGTTTT	
TACATCCTTA	
GCACCAAGCTT	
CTTCCAACAG	
TTTCTTAACAA	

236		
CGATTTATAG TTGCTCCTGT	AGTATAGATA TCATCTATAA GTAGGATTTT TTTAGGAATA	4620
GTGACTCCAC TTTTAATAAA	GAAAGGAAGT TCTGCCCCA AGCGCTCTGA ACGATTTTA	4680
GAAGAACTGG CTCTCTCTTC	TCTTTCTCT AATAAATCCA GATACTCAA GCCTGCTGCC	4740
TCTACCAAGC CCTCAACCTG	ATTAATCCT CTATTAGCAT ATCTATCAGG ACTTAGGGGA	4800
ATTACAAACAA ATTGATACTC	TTTGTACTTT TTCAACTCCT CACTTAAAAA TGAAGCGAAA	4860
ACTTTCTTA ACAGGAAGTC	TCCATCAAAC TTATACCGAC TGAAAAAATC CTTCATAGCT	4920
TGATTGTAAG TAAAATCGC	TCTATGACTG ACTTCAACTC CCTCTTTACA CCAAAGTTGA	4980
CAATCTGAC ACTTTGTTGA	CAACTCTGTT TTCATACAAT TTGGACAGTT CTCTTCCCCA	5040
ATTCTTCAA AAGTAGAATC	ACAGTCTGAA CAAAGACAAG AGTCATCATT CCTCAGAAGT	5100
AAGAGACTAC TAAAAGTTAA	AACAGTCTTC ATAGTCTGCC CACATAACAA GCACTTCATA	5160
GACCAGCCTC CTTATTCAATC	ATCTGAATTT CCTTAATCGC CTTCTTGATT GAAGCATTAA	5220
ACCCATCATG GAAGAAAAGC	AAATCTCCCG TCAGCTCTAC CATGCTTCGT CCAACTCGTC	5280
CACCAATCTG AATCAAACTA	GACTTGGTAA ACAACGATG ATTGGCCTCT ACTACGAAA	5340
CATCCACACA AGGAAAGGTA	ACTCCGCGCT CCAAGATTGT CGTACTGATA AGTATTGTCA	5400
GTTCTCCATC TCGAAAAGCT	TGTACTTGCT CTAATCGATC CTCTGTTACA GAAGATACAA	5460
AGCCAATTTC CTCATTTGGA	AATTGCTCCT GTAAAGATTTC TGCTAACTGC TCCCCTTCT	5520
TAATTCTGA AGCAAAATG	AGTAACGGAT AAGCTGTCTT TCTCTGCTTC TCAATATAGG	5580
ACTTTAACCTT TGGTACAAA	CGATTCTTGT CTAAGTAGCG ATTAAAATCC GATAACCAA	5640
TTGGTTTTGG AATAATCAAC	GGATTCCAT GAAACCGTCT CGGTAAATTG AGTCTTTTA	5700
GTTCTCCTAA ACGGACCTTT	TTATCTAACT CATTGGTCGA AGTCGCTGTT AAAAGATTC	5760
TCAATCCATT CTCCCTTACA	CTATTCTTGA CAGCGTGGTA AAGCATGGGA TTATCAACAT	5820
AAGGAAAAGC ATCTACTTCA	TCCACTATCA GCAAATCAA AGCTTGATAA AACTTCAATA	5880
ACTGATGGGT TGTGCAACA	ACTAGTGGTG TTGAAAATA AGGTTCCGAT TCTCCATGTA	5940
GCAAAGCTAT CCCCAAGAA	AAATCCTGTT GCAGGCGCTT GTACAGCTCC AAACAAACAT	6000
CTATGCGAGG ACTAGCCAA	CACACTGCAC CACCCGCATT GATCACTTTA GCCACTACTT	6060
GATAAATCAT TTCTGTCTTT	CCAGCTCCTG TTACCGCATG AACTAAGGTT GGCTTTGCT	6120
TGTCTACTAC TTGAAGCAAT	CCCTCTGACA CCTTCTCTTG AAAAGGAGTT AATTGGCCGC	6180
GCCATTTGAG AACATCTTGC	TTTGGAAAAT CCTCCTGCCG AAAATAGTAT AAAGTTTGAT	6240
CACTTCTGAC TCGCTTCATC	AGCAAGCACT CTCGACAATA GTAAGCACCG ATGGGCAAAT	6300
ACCATTCTTC TAGAATAGTA	CTATTACAGC GTTGACAGAA AAGTTCCCC TTCTCCTTTC	6360

TCATTGCTGG AAGTTTCTCC GCCAACTGAC GTTCTTCTTC TGTAAATTCA TTCTCAGTAA	6420
ATAAACGACC GAGATAATCT AAATTTACTT TCATACTTCT TTATTCGTA AAACTAGCAC	6480
TTTAGATGAT TTTTAGTAC AATTAATCA TGGAAATTAG GACAATTAAA GAGGACGGTC	6540
AAGTCCAAGA AGAAATCAA AAATCTCGCT TTATCTGCCA TGCCAAGCGT GTTTATAGCG	6600
AAGAAAGAGGC TCGTGAACCTTC ATTACTGCCA TCAAAAAAGA ACACATACAA GGCACACATA	6660
ACTGCTCTGC CTTCATTATT GGAGAACGTA GTGAAATTAA ACGTACAAGT GATGATGGTG	6720
AGCCTAGTGG TACTGCTGGT GTTCCCATGC TTGGGGTACT AGAAAATCAC AATCTCACCA	6780
ATGTCTGTGT GGTCGTGACA CGCTACTTTG GTGGTATTAA ACTAGGCCT GGAGGACTAA	6840
TTCGTGCTTA CGCCGGCAGT GTGCCCTTAG CTGTCAAAGA AATTGGTATT ATTGAAATAA	6900
AAGAACAGGC TGGCATTGCT ATTCAAATGT CTTATGCTCA GTACCAAGAG TACAGTAAC	6960
TCCTTAAAGA ACATGGCTC ATGGAGCTGG ATACAAACTT TACAGATCAA GTCGATACGA	7020
TGATTTATGT TGATAAAAGAA GAAAAAGAAA CTATTAAGC TGCACTTGTG GAGTTTTTA	7080
ATGGAAAAGT CACTTTAAGT GACCAAGGTT TACGAGAGGT TGAAGTTCTT GTAAACTTAG	7140
TCTAAACAT GAATAATACA GCGTTTCGTT GACATTCTCA CAACTACTTT AGCGAGCAA	7200
ATAAAAAGAG GCGTACCAAA ATATACTAGA AAATGAAGCA ATTCAAACGA AACCTGATAT	7260
CGTTTCCCTT CACACCTATT TACTAGAATT AGCTGAACCC AATCACTTGA AAATTAATGA	7320
CTTTGATCTA TGATATATAG AAATGGTATG GATAGCGTTA TACTAAACAT ATCTTATACA	7380
AAGAGGTATT CATATGTCTA TTTATAACAA CATTACTGAA TTAATCGGTC AAACACCGAT	7440
TGTTAAACCTT AACAACATCG TGCCAGAAGG TGCTGCAGAC GTCTATATAA AGCTTGAAGC	7500
ATTTAACCTT GGTCATCTG TAAAAGACCG TATTGCCCTT AGCATGATTG AAAAGCTGA	7560
ACAAGATGGT ATTCTGAAAC CTGGTTCTAC TATTGTTGAA GCAACAAGTG GAAACACCGG	7620
TATTGGACTT TCATGGGTAG GTGCTGCTAA AGGGTATAAA GTCGTCATCG TTATGCCCTGA	7680
AACTATGAGT GTAGAACGAC GTAAAATTAT CCAAGCTTAT GGTGCTGAAC TCGCCTAAC	7740
TCCTGGTAGC GAGGAATGAA AAGGTGCTAT TGCTAAGGCT CAAGAAATCG CTGCTGAACG	7800
TGATGGTTTC CTTCTCTTC AATTGACAA TCCAGCTAAT CCAGAAAGTAC ACGAAAGAAC	7860
AACAGGAGCT GAGATACTAG CTGCTTTCGG TAAAGATGGA TTAGATGCCT TTGTTGCTGG	7920
AGTAGGTACT GGTGGAACGA TTCTGGTGT TTCTCATGCA CTCAAATCG AAAATTCTAA	7980
CATTCAAGTT TTGCAAGTAG AAGCAGATGA ATCTGCTATT CTATCTGGTG AAAACCTGG	8040
TCCTCACAAA ATTCAAGGTA TCTCAGCTGG ATTTATTCCCT GATACACTTG ATACTAAAGC	8100

238

CTATGATGGT ATCGTTCGTG TAACATCAGA TGACGCTCTT	GAACTCGGAC	GTGAAATTGG	8160			
TGGAAAAGAA GGCTTCCTTG TAGGGATTTC	CTCAGCTGCA	GCTATCTACG	GAGCCATCGA	8220		
GGTTGCCAAA AAATTAGGTA CAGGTAAAAA	AGTCCTTGCC	CTAGCACCAG	ATAACGGTGA	8280		
ACGTTATCTC TCTACAGCAC	TTTATGAATT	GTAACCGTCC	AATAACGAAG	TCTATTGAAA	8340	
AATCTCCAGA CTAGAGAACT	CACGGATAGT	TCCTAATCTG	GAGATTCTT	ATTTGCACTT	8400	
TTCTTGACCA	ACTTTAGTCC	ATGGTAAATA	GGCCTCTAAA	ACCTCTTTGT	TTACGAGAGT	8460
TTCCACGTTT GGAAGACATT	CTAGAAAGATA	GGATAGATAT	TTCTCACTAT	TTATAATGGA	8520	
TTGAAATAAG ATATGACAA	ATCGATTAGA	ACATGATGGT	AAAGCGTAAT	CCCTTGTTTC	8580	
TCAGCTTTCC CAGACAAAAA	AGTCCAATAG	TAAGTCAGCT	GAATATCACT	CTCTAGCACC	8640	
CTATAAGAAG TTTCATCCGC	ATGAAGTAAG	GGCTGAGTC	ATAGTCTCTC	TCGCAAGAGG	8700	
TTATAAAGGG GCTCCAATA	GTATTGACTC	GTCTTGATAT	GCCAATTAGA	GATTTCCCTA	8760	
CGTGTGATTG GTAAACCCAT	CCTAGCCAA	TCTTCTTCTT	GGCGATAATT	GGGTACCTTC	8820	
AGATTAAACT TCTGATGGAT	GGTGTGAGCG	ATAATAGAAG	CTGAGCCAAA	GTTATGCGCT	8880	
AAAGGGCTT TAGGAATAGG	AGCTTTCACA	AGCTTATCCA	GATGATTATC	TTTTACTCGT	8940	
TATGGACAAT GCTATATGGC	ATAATCAAG	TACCTTAAAG	ATTCCGACTA	ATATTGGCTT	9000	
TGCATTATT CCTCCATACA	CACCAAGAGAT	GAACCCCAT	GAACAAAGTGT	GGAAAGAGAT	9060	
TCGTAAACGT GGATTTAAGA	ATAAAGCCTT	TCGAACTTTG	GAAGATGTCA	TACAAGGACT	9120	
GGAGAAGGAG GTGATAAAAGT	CCATCGTTAA	TCGGAGACGG	ACTAGAATGC	TTTTTGAAAAA	9180	
CAGATGAGTA TAAAAAGAAA	GTCCCTCATTT	CAATAGAAAT	CACGACTTTC	TGATGAATT	9240	
ATAGTAAAT GAAATAAGAA	CAGGATAGTC	AAATCGATTT	CTAACAAATGT	TTTAGAAGCA	9300	
GAGGTGTACT, ATTCTAGTTT	AAATCCACTA	TATTTGGGA	GTGATAGAAA	AGCCCTTCAT	9360	
CAGCCAATCT	ACTTGTTCA	GTGCGAGAGC	TTTGACATCC	TTTTCTGTAC	TGGACCAAGT	9420
CAGTTTCCG TTCTCAAAGC	GTTTATATAA	TATCCAAAT	CCTTGACCAT	CCCAGTAAAG	9480	
AACTTTAAAG CGGTCTTAC	GTCCACCACA	AAAGAGAAAG	ACTTGATCGG	AGAAAGGATC	9540	
CAATTCAAAG TGGTTTAA	CTACATAGGC	TAATGAGTCT	ATTCCCTGCC	TCATATCTGT	9600	
CTTGCCACAA ACAAGGTGAA	CTTGACCTAA	ATCACTTAGT	TGAATTATCA	TAGTACAATA	9660	
CCTTTCTCC GATAATTATT	TTTTATCTGG	TATACTGGAA	GTTGGGGAAAT	TAGGATAGAT	9720	
ACCTTGTTAT GACCGCCTTA	CTATGAATT	GAAGTATAGT	CTCCTAAATG	CACTTAGCCC	9780	
TTATTATAGG GCTTTTGTT	TTAATTATTC	TAATCGAGTG	AGACTGGGA	AAAAACAATT	9840	
TCAGGAAAAA TCTAAGCCCT	ATACAAAAAA	GGAAGCAATT	TGCTTCCTTT	CTATTATTAG	9900	

TTATTCAAGG CTGCTGCCAT TGTAGCTGCA ACTTCAGCTT CGAAGTCGTT TGCAGCTTTC	9960
TCGATACCTT CACCAACTTC AAAGCGAGCA AACTCAACTA CCGAAGCGTT AACTGATTCA	10020
AGGTATGCTT CAACTGTCTT GCTGTCTGCC ATGATGTAAA CTTGTGCAAG AAGTGTGTAA	10080
GCTTGGTCAA CTTTAGTGTGTT ATCAAGCATG AAGCGATCCA TTTTACCTGG AATAATTTG	10140
TCCCCAGATTT TTTCTGGTTT GCCTTCTGCA GCCAATTAG CTTTGATGTC AGCTTCAGCT	10200
TGAGCAATAA CATCATCAGT TAATTGAGCT TTTGATCCAT ACTTCAAGTG TGGAAGAGCT	10260
GGTTTATTAA CCATTGCACG GCTTTCGTTG TCTTGGTCGA TAACGTGATT CAATTGTGCC	10320
AACTCATCTT TAACGAATTG CTCATCCAAT TCTTGTAAAG AAAGAACTGT TGTTTCATC	10380
GCTGCGATGTC GCATTGACAA TTGTTTAGCA AGTGCTTCGT CTCCACCTTC AACAACTGAA	10440
ATAAACACCGA TACGTCCACC GTTATGTTGG TATGCTCAA AGTGTGTC GTCTGTTTTT	10500
TCAATCAATG CAAAGCGACG GAATGAGATT TTCTCTCCGA TAGTTGCTGT TGCAGATAAG	10560
TATGCAGCTT CAAGAGTTTC ACCTGAAGGC ATTATCAAAG CAAGAGCTTC TTCGTTGTTA	10620
GCAGGTTTC CTTCAGCAAT GACTTTAGCT GTAGTATTTA CCAATTCAAC GAATTGAGCG	10680
TTTTTTGCAA CGAAGTCAGT TTCAATCGCT TTTCGATGT CACCGCTCTGT TTCTACAAGC	10740
ACATAAACAC CAGTCAAACC TTCTGCAGCA ACACGGTCAG CTTCTTAGC TGCCCTAGCC	10800
ATACCTTTT CACGAAGCAA TTCAATCGCT TTTCGATGT CACCGCTCTGT TTCTACAAGC	10860
GCTTTTTAG CGTCCATAAC ACCGGCACCAC GATTTTTAC GCAACTCTTT TACAACTTTA	10920
GCTGTAAATT CTGCCATTAA ATTCTCCTA TATTTTTGAA AAATAGGAGA GCGCGGCTAA	10980
CCCCGGCCTC CGG	10993

(2) INFORMATION FOR SEQ ID NO: 16:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 8411 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 16:

CGACGGGGAG GTTTGGCACC TCGATGTCGG CTCGTCGCAT CCTGGGGCTG TACTCGGTCC	60
CAAGGGTTGG GCTGTTGCC CATTAAAGCG GCACGCGAGC TGGGTTCAGA ACCTCGTGAG	120
ACAGTTCGGT CCCTATCCGT CGCGGGCGTA GGAAATTGAGA GAGGATCTGC TCCTAGTACG	180
AGAGGACCAG AGTGGACTTA CCGCTGGTGT ACCAGTTGTC TTGCCAAAGG CATCGCTGGG	240

240						
TAGCTATGTA	GGGAAGGGAT	AAACGCTGAA	AGCATCTAAG	TGTGAAACCC	ACCTCAAGAT	300
GAGATTCCC	ATGATTATAT	ATCGATAAGA	GCCCTGAGAG	ATGATCAGGT	AGATAGGTTA	360
GAAGTGGAAAG	TGTGGCAGACA	CATGTAGCGG	ACTAATACTA	ATAGCTCGAG	GACTTATCCA	420
AAGTAACGTGAA	GAATATGAAA	GCGAACGGTT	TTCTTAAATT	GAATAGATAT	TCAATTGTA	480
GTAGGTATTA	CTCAGAGTTA	AGTGACGATA	GCCTAGGAGA	TACACCTGTA	CCCATGCCGA	540
ACACAGAAAGT	TAAGCCTAG	AACGCCGAA	GTAGTTGGGG	TTGCCCCCT	GTGAGATAGG	600
GAAGTCGCTT	AGCTTTAAC	CGCCATAGCT	CAGTTGGTAG	TAGCGCATGA	CTGTTAAC	660
TGATGTCGTA	GGTCGAGTC	CTACTGGCGG	AGTAATTGAT	AAAAGGGaAC	ACAGCTGTGT	720
TCCTCTTTT	GTATCAATT	GTATCACCAA	GCATTTCAT	AAGGAAGTCT	GTTATTTCTT	780
GAGAACTTTC	TTTTTTCCA	TGTGCAATCC	AAGTTTGGCA	GACACAAAAA	AGTGCATGAG	840
TTAGATAGAT	GCTACTATAT	TCTAATTCA	TGGTATTAG	ATTCAGTTGC	ATAAATCGCT	900
TTTGATAATC	TGTACTAACG	ATGATATGAA	GTTTATTCG	TAAGAAATT	TGGATTTCTT	960
TAGTCCCATT	TTCAGAAAGA	AGGCAGCCA	GAAGTGGTC	TGACTCTAGA	TATTCAAAAA	1020
CTTCTAAAAT	AGCGCTCTT	TTGTGATGAG	CATGTTTG	AAAAATATAT	TCAAATGTAT	1080
GGAATAGCTT	GCTTGATAG	TGCTCAATCA	TATCATACTT	ATCCTTATAG	TGAGTATAGA	1140
AGCTGGAACG	ACTAATTCCG	GCTTTCTA	CTAATTGAC	AGTAGAAATT	TTATCAAATG	1200
GCTGTTCCAT	CAGTAATTGT	ACCATAGCAT	TTCAATAGT	TCGTTTGTT	TTAAGCGTT	1260
TGTTACTTTC	TTGCATATTT	CCTCCTGTA	AACAAATTAG	ACTATATGTC	AAAAATAGA	1320
TTTTTTATCT	TGTAATTAG	ATTTTTAAT	GTATAATCTA	TTATATCAA	ATTTTAGACA	1380
ATATGTTAA	AAAAGGAGAA	ACTAAGTTA	AAGAATGGAA	AGCAATTAA	AAAAAACCAA	1440
CCTTTATTAT	TGTCAATGATC	GGGATTTCTC	TTATCCAGA	TCTGTACAAT	ATCATATT	1500
TGTCATCAAT	GTGGGATCCA	TATGGCAAT	TGTCTGACTT	ACCTGTGGCA	GTTGAAATA	1560
ATGATAAAAGA	GGCTTCCAT	AATGGTAATA	CTATGGCAAT	AGGAAAGAC	ATGGTGTCCA	1620
ATTTAAAAGA	AAATAAAACC	TTGGATTTTC	ATTTGTAGA	TGAAGGGAA	GGAAAGAAGG	1680
GATTGGAAGA	TGGCGATTAC	TATATGGTAG	TGACTTTACC	AAGTGAATT	TCTGAAAAAA	1740
CAACTACATT	ATCCAATATT	CAATCGACAG	CAGCTTATCA	ATCATTGACA	AGTGAGCAAC	1800
AAACTGAGAT	AACTGATTCT	GTATCTCAA	ATTCAACTGA	TAGTATTCAA	TCGGCTCAGT	1860
CAATTGAGC	TTTAGTACAA	GATTACAGG	GAAGTTAGA	AAACCTACAA	AATCAATCTT	1920
CTAATCTTTC	GACTTTAAA	AATCAATCTA	ATCAAGTATC	ACCTATTACT	TCTACTTCTT	1980
TGATAGGATT	GTCAAGTGG	TTAACAGAGA	TACAAGGAGA	TGTTACTAGC	AAATTAGTTC	2040

241

CTGCCAGTCA	GTCGATTGCA	TCAGGTGTA	ACGCATATAC	TACAGGTGTT	GATAAAGTTT	2100
CTCAGGGCGC	AAGTCAACTA	AGTGAAAAAA	ATGCCACCTT	GACAGGTAGT	TTGGATAAAC	2160
TAGTTTCAGG	CTCAACACC	TTGACACAAA	AATCTCTAG	ATTGACAGCA	GGAGTTGGTT	2220
AATTACAATC	AGGATCTGGG	CAATTAGCAG	ACAAATCCAG	TCAGTTACTT	TCAGGTGCTT	2280
CTCCATTAGA	GAATAGAGCT	AATAAATTGG	CAGATGGATC	TGGGAAACTA	GCAGAAGGTG	2340
GAACAAAGTT	AACTCTGGA	TTGGAAGATT	TACAGACAGG	ACTTGCTTCT	TTAGGACAAG	2400
GACTAGTAA	TGCTAGTGAT	CAACTCAAAT	CACTATCAAC	AGAATCTAA	AATGCAGAGA	2460
TTTTGTCAAA	TCCACTCAAT	CTTCAAAAAA	CAGACAATGA	TCAAGTCCT	GTAAATGGAA	2520
TCGCAATAGC	TCCTTATATG	ATATCAGTTG	CTCTTTTTT	GCAGCAATAT	CAACAAATAT	2580
GATATTGCG	AAATTGCCTT	CAGGACGTCA	TCCAGAGAGC	CGTTGGGCTT	GGTTGAAATC	2640
TTGAGCTGAA	ATAAATGGTA	TTATAGCTGT	TTTGGCAGGA	ATTTTGGTAT	ATGGAGGAGT	2700
TCAGCTTATT	GGTTAACTG	CTAATCATGA	GATGAGAATA	TTTATTCTCA	TCATCCTAAC	2760
AAGTTTAGTA	TTCATGTCTA	TGGTGACCA	TTTAGCAACG	TGGAATAGCC	GTATAGGAGC	2820
TTTTTCTCA	CTTATTTGC	TTTTACTACA	GTTAGCATCA	AGTGCAGGTA	CTTATCCACT	2880
TGCTTGTACA	AATGATTTCT	TTAGATCTAT	TAATCCCTGG	TTACCAATGA	GCTATTCTAGT	2940
TTCGGGATTA	CGACAAACAA	TCTCTATCAA	CAAGTCATTT	TCCTAGCTGT	CATACTAGTT	3000
CTATTTACTA	GTTTAGGTAT	GCTAGCCTAT	CAACATAAGA	AAATGGAAGA	AGATTAAGAA	3060
AATCGACCGA	TTAACTGGTC	GATTTTTAT	GCCTTAGATG	ACTTCGCT	GTGATTATAG	3120
ATTCCAAATA	GTAAGAGAGA	AGTAAAGGAA	CAGATTGCTC	CGATAATAAA	ACCATTGGGA	3180
ATGAAGGAA	GTGTAATAGT	TCCTTCCCC	TTGGGAATGT	CAACTTCAT	AAATCCAGTT	3240
TGAGCTGTT	TAATTTCTAT	TTCTTACCA	TCTTGGTAGG	CAGACCAACC	TTTGTCTAA	3300
GGAATGGTGA	AGAAAATAGA	TGTATCTGT	TGGACATCAT	ATGTAGCAA	AACCTTGTT	3360
TTAGAAAGTTG	ATACTGTGAC	AGGTGTTCT	TTAATTTTT	GAATTGCTC	GGTGAAGTT	3420
TTGGTATCTA	AACGATAGAA	GGTAGGAGAT	TCAAATGATA	CTTGTGAATT	TCCAGGGAAA	3480
CTAACATTGA	TATTGAAAGT	TTTTTCTCT	TTAGTATATC	CTAGATTAAA	GAAGGAGAAG	3540
ACATTATCAG	TTGTAAAAGT	CTTTTTTCA	CCATTACAA	GGATGTCAAC	CTTCTTTGT	3600
TTATCGTTAG	AAAATGAAG	GTTTATGAAA	GAGAGATAAA	CTTGGCTGTT	TTCTGGAAC	3660
TCAATTGAT	ACTGGATTGC	TGCATCTCA	TTTGAAGAAC	TTGTGACACT	AATCAAATCA	3720
TTAGTATTTT	CTATTTTTC	TGTTTTTCA	TAAGGTATTG	GAGAAAATA	ATCAAATTC	3780

242							
ACGTTAGCAA	GTTGATTTAA	AAATGAGGCC	TGATTATCCA	AGGTATGTT	ATTGAAC	TTG	3840
ACATCATTGT	AAACAGATTG	ACTCGCAACT	GCAATCGGAA	GAGAGTATTG	ATTTTCATAT		3900
AGGGTAAGAT	TATCTTTTG	ATAGATATCT	TTAAAGCCAT	ACTTATCAAT	AGGACTGTCT		3960
GAGATATTGT	ACTGGATACC	AAATAAACTA	TCAGCCAAA	TACTATTATT	TGCATATCGG		4020
AGATTGAGAT	TAGTCCCAGA	GGATTTAAA	CCAAGTTTAT	CTAAAGTAGA	GCTTGATGAA		4080
CGATTTGAA	CAGATGAAA	TTGAGAGATT	CCATTGAGT	TGAATTTCAT	ACTGTCATT		4140
CCTGTCAG	TTTGAGTTT	TTCAGTACGA	GTAAATTGAT	TTCCAATATA	TGTTGAGAAA		4200
GATTCCATAG	CTGGGATATC	TCGACTATAA	GCACCTCGAG	AAGCAAATCC	CCATTCCCTTA		4260
GCAATTCCGT	CCATTGAGA	TGAAGCATT	AAACTCATT	CAACCAAGTAT	AAATAAAGAG		4320
ATTAGAATGG	CAAATAGATT	CACAGATATA	AACTTTTG	TAACTGCAAG	GAGTAAAAGA		4380
GAATAGACAA	CCAAAATTC	AAGAGTAAGC	AGAATATTCA	AATCTGTTAA	AAAAGAATAA		4440
TGCGATTTA	GATAGATGGT	AGCTAAAAT	CCTGCTACTA	CAAGAAAAAG	CGAAACTAAA		4500
AAATTCCAGA	CTTTAAGTTC	TTTCAGACGC	TTTAAGACTT	CTGCTGCTGT	GTAAATTAAAC		4560
AAGGTAGAGA	AAATCCAAGC	ATAGCGATGT	AAAAACATGT	TTGGAGTATG	CATGCCTTGC		4620
CAAAATAAGT	CAAGAGCTTC	TATGTAAG	CTTGCAATTA	GAATGCAA	GAATATTACA		4680
TATATGAGTT	TCACGTGAA	CTTAATAGAT	TTCAAGCTAA	AAAATAAAAT	GGTCAAAATA		4740
AAGGGAAATA	GTCCAACAAA	AATCATGGG	ATGGCCCCAT	ACTTTGTTGT	GTCAAAGGAA		4800
CCAATGAATT	GCTTAGAAA	GAGATCAAGA	TACCAAGTAC	TTTCAGTTTG	AAACTTTGTA		4860
ACTTCAGTCA	ATTTTCCCC	ATGTGTCTGT	AAATCAAATA	GAGTGGGAAG	AQTCATAATC		4920
AAACTAGCCA	TACCAAGTAA	AAAGGAGATA	ACTATGAAAT	CAAGAACAGA	TGATTTTCGA		4980
GTCTTAAAGT	CCCACGAAAT	TTGACAGAGA	TACCAAGAAA	TAAGAAACAA	TACTGTCATA		5040
TATCCAAAAT	AATAATTTG	AATAAATAAG	ATTGACAGAC	TTGTAAAGTA	CAATAGGAGT		5100
TTCTTTTCAG	TTATCAGTAG	ATGTAAACCA	GTTATAATTA	AAGGAATCAA	GATAAAAACA		5160
TCTAGCCAGG	TTTTTATCTC	TAATTGACTG	ACAGTGAAAC	TCATCAGAGC	ATAGGAAGTA		5220
GATAAGGCTA	GTTTAAAAT	CTGAGGGATA	GATTGAAACA	ATTTATTCAA	ACTAAAAAAG		5280
GTTGACAGAC	CAATCAATCC	AAATTTTAAG	AGAGTTGTCA	GATAGATGC	ATCTGGCATA		5340
TTCGTTAGAT	CAAAAAAGTA	AACCAGAGGC	GGCAGAAAAC	TACCCAAGTA	ATAACTAGAT		5400
AGGGCATAGA	AGTTTAGCCC	TAGACCACTT	GTAAAGGTGT	AAAACAGATT	ACTATTTC	CCA	5460
TGTAGGATAT	TTCGTAAGGC	TACATCAAA	ATAACGTATT	GATGAAAGCC	ATCTCCTAAT		5520
AGAGGAGAGT	TGTCGCTATT	CCAGTAGATA	CTTGAGATA	GATATACTCC	AGACATAATC		5580

ACTACAGGAA	TGATGAAAGA	AATAAAATAG	GTTCGATATG	TTTTTAAAAAA	TGATTCATG	5640
TTACCTCGTA	GAATGATAGA	AAAATCAGTT	GGTTAACCCA	ACTGAGTTTT	GAAGTTTAT	5700
TTAGCTTTC	CAAAGTTCTT	TAACTTTGC	TTGTACTTCT	GCATTTCTA	GGAATTCATC	5760
GTAGGTTCA	TCGATACGGT	CAATGACGCC	ATTTTAGAT	AAGACAATGA	TATGGTTAGC	5820
CAAAGTTGA	ATAAATTCTGT	GGTCATGGCT	GGCAAAGATG	ATTGATTCTT	AAAAGTTTT	5880
CAATCCATCA	TTCAAGCTTG	AGATAGATTC	CAAGTCCAAG	TGATTTGTTG	GATCATCAAG	5940
TACAAGGACA	TTTGATTTTA	AGAGCATGAG	TTTGAAAGC	ATGACACGAA	CTTTTCTCC	6000
CCCTGACAAG	ACATTTACAG	GTGGTTAAC	TTCATCTCCA	GAGAAGAGCA	TACGGCCGAG	6060
GAAGCCACGT	AGGAAAGTAT	TGTCATCTTC	TTCTTTACTT	GCAGATTGAC	GCAACCAGTC	6120
AAGAATTGAT	TCTCCTCCTG	CAAATCAGC	TGAGTTATCT	TTTGGTAGGT	AAGATTGACT	6180
AGTTGTAAC	CCCCACTTGA	CAGTTCCCTTC	ATAGTCATA	TCTCCCATGA	TTGCACGAAT	6240
TAATGCAGTC	GTGGAAATAT	CATTTGTCC	AATAAGTGCT	GTCTTATCAT	CTGGACGC	6300
GATGAAACTA	ATATTATCCA	AGATAGTTTC	ACCATCAATC	TTTACAGTTA	AAATTCTAC	6360
TGTCAGAGA	TCATTACCAA	TCTCACGTT	CGCTTTAAAG	TTGATAAAATG	GATATTACG	6420
ACTAGATGGC	ACAATCTCTT	CTAGCTCAAT	CTTATCAAGC	ATTCTCTTAC	GTGATGTTGC	6480
CTGCCTTGAC	TTAGAAGCAT	TGGCAGAGAA	ACGAGCAACA	AATTCTTGCA	ATTGTTTAAT	6540
TTTTTCTTCT	GCCTTACCAT	TACCGCTCTCC	TAGCAATTAA	GCAGCAAGCT	CAGAAGATTG	6600
CTTCCAGAAG	TCGAGTTTC	CGACATAGAG	TTTGATTTTT	CCAAAGTCAA	GGTCGGCCAT	6660
GTGAGTACAA	ACTTTGTTA	AGAAGTGACG	GTCGTGGAT	ACTACGATAA	CTGTGTTATC	6720
AAAGTCAAATC	AAAGAAGTCTT	CTAACCAAGT	ATACGATTGG	ATATCCAAAC	CGTTAGTAGG	6780
CTCGTCCAAG	AGAGAACAT	CTGGTTTAC	AAAAAGTGT	TTGGCCAGGA	GAACCTTTAC	6840
TTTTTCACCG	TTGGCCAATT	CGCTCATGTT	TTGGTAGTGT	AATTCTCTG	GAATGTTAG	6900
GTGTTGAAGT	AGTTGAGAGG	CTTCACCTCTC	TGCTTCCCAA	CCTCCAAGTT	CGGCAAAC	6960
TCCCTCGAGT	TCGGCAGCAC	GAACCCCGTC	CTCGTCTGAG	AAATCTTCT	TCATGTAGAT	7020
AGCATCTTTC	TCTTCATGA	TGCTATAAAG	TTTTTCATTT	CCCATGATAA	CGACATCAAT	7080
GGCACGTTCA	TCTCGTAGT	CAAAGTGATT	TTGACGAAGA	ACAGAGAGAC	GTTCATCTGG	7140
ACCAAGAGAG	ATGTGACCAAG	TAGTAGGTT	GATATCTCCA	GCTAAAATTT	TTAAAAAGGT	7200
TGATTTCCG	GCACCAATTAG	CACCGATTAA	TCCGTAAGTA	TTTCCCTCTG	AAATTGAT	7260
ATTGACATCA	TCAAAAGTT	TGGCATTCACT	AAAACGTAGT	GAAACATCAG	ATACTGTAAG	7320

244

CAATGTTTT CTCCTATATG TGTAATATAT TTATTCTACT AGAAAATACA GAAATATTCA	7380
AATTTTATT TGTCAATTT GTGTAATTA TATTTACAGT ATCCTTACA CAAATCTGTA	7440
AAAAGCAAGG CTGATTTATT TTGATAAATT ACGGTTATTT CATTAAAAAA ATGCTATAAT	7500
TGAAAGGACT ATATCGAAGG AGAACAAAAT GACTAAACCC ATTATTTAA CAGGAGACCG	7560
TCCAACAGGA AAATTGCATA TTGGACATTA TGTTGGAAGT CTCAAAATC GAGTATTATT	7620
ACAGGAAGAG GATAAGTATG ATATGTTGT GTTCTTGGCT GACCAACAAG CCTTGACAGA	7680
TCATGCCAAA GATCCTCAA CCATTGTAGA GTCTATCGGA AATGTGGCTT TGGATTATCT	7740
TGCAGTTGGA TTGGATCCAA ATAAGTCAAC TATTTTATT CAAAGCCAGA TTCCAGAGTT	7800
GGCTGAGTTG TCTATGTATT ATATGAATCT AGTTTCGTTA GCACGTTGG AGCGAAATCC	7860
AACAGTCAG ACAGAGATT CTCAGAAAGG ATTTGGAGAA AGCATTCCGA CAGGATTCTT	7920
GGTCTATCCA ATCGCTCAAG CAGCTGATAT CACAGCTTTC AAGGCTAATT ATGTTCTGT	7980
TGGGACAGAT CAGAAACCAA TGATTGAGCA AACTCGTGAA ATTGTTCGTT CTTTTAACAA	8040
TCCATATAAC TGTGATGTCT TGGTAGAGCC GGAAGGTATT TATCCAGAAA ATGAGAGAGC	8100
AGGGCGTTTG CCTGGTTTAG ATGAAATGC TAAAATGTCT AAATCACTAA ATAATGGTAT	8160
TTATTTAGCT GATGATGCGG ATACTTTCG TAAAAAAAGTA ATGAGTATGT ATACAGATCC	8220
AGATCATATC CGCGTTGAGG ATCCAGGTAA GATTGAGGGAA ATATGGTTT TCCATTATCT	8280
AGATGTTTT GGTGTCAG AAGATGCTCA AGAAATTGCT GATATGAAAG AACGTTATCA	8340
ACGAGGTGGT CTTGGTGATG TGAAGACCAA GCGTTATCTA CTTGAAATAT TAGAACGTGA	8400
ACTGGGTCCG G	8411

(2) INFORMATION FOR SEQ ID NO: 17:

- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 9064 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 17:

TGCCGTACTC AAGTACAGCC TGCCTAAGT TTCTAGTTT GCTCTTGAT TTTCATTGAG	60
TATTAAGTAAAC CAAAATCCGA CCACATAGCC AGCCCTATG AATATAGCCA TTAAAGCTAG	120
CATGGAATTT AGGAAATTAA AAACCACCGC AGATACAAAG GTTAGCACAA AAACATTAAG	180
AGCAATGGTG TCAGAAGCCA AGACTAGAAT ATAGGGTGTC AACCGATCTA AAGTTTGGA	240
ATCTAGGAAA ATAAGTGTGTT TATACATGAT GACCTCCTCT ATGGCTGAAA AGCAAGCCTT	300

245

TTGTTTTTTT	ACCCCAAGAC	CCTATGTAGA	AAAGTGAGCA	AAAACGGAA	GGTCGCTACA	360
ATATTATTGA	TCACATGCAC	CGCATAGGAT	GGATAAATGC	TCTTGGTATA	GCGGGTCAAA	420
CCAGCAAAGA	TGATTCAC	TGTTGCAAAG	ACGAAGATAT	CTAACAGACT	AGGCAGGCTT	480
GAAAAATGAG	GGAGAGCAAA	TAAAATAGAA	GGAAGAAGCA	AATCAAGACC	AAATCGCGAA	540
TGCTTAAAGA	AAGCATGTTG	CAGTAATCCT	CTATAAAATCA	ATTCTTCCAT	CAGTGGAACCC	600
AGAAAGAACAA	GGGCTATATA	AATACCTAGC	TCTGCAAAGT	TAGTCCCCT	ATAACCAATC	660
AATACAGCCC	AACCTTCCGC	AGTTGACTGA	ACATGTTAG	CTGCTGAAC	GTTAAAAGAG	720
ATCTGGAACA	CTAGCACTAA	TACTGTCAA	ATCGAATACC	AAAGCCATT	TTTTCTTGGAA	780
ATGCGGAAGA	GATAACCATG	GCCTGTCTTA	ACAAGAACCA	CAATCATGAC	TCCAATAAAA	840
AGTAAACTCA	AGATATTTG	AATCCAGAAT	AAATTGCCTA	TCTGAGAAGA	AAATTGCCAA	900
TAGTTTGGA	CGATAAGCGT	CAGCTGAGAA	AGACTAAATA	CGAAAAATAA	GTAAGAGAAG	960
ACTGCACTTA	TTTGAAATAG	AAGTTGATAC	TTTTTCATAG	AAATCCTCCC	TACTATGACC	1020
TCACCTTGTC	AGGCTCTACT	GCTGTAAGAT	TAAGAAGACA	GTTTGT	TTTAAGGCTA	1080
ACCTGACTAC	TAGATAATAG	ATACATTAAG	GCATTAAAGA	CAATGAAAAT	ATGTCCATAG	1140
AATAAAATCA	ACCTCGCATC	CAAACCAAGA	TAAAGTTGA	TTATCAAAAA	GATGAGCAAA	1200
AGAATTGAA	ACCATAAGGT	TTTCCAAAA	ATAAATTAA	AGCGATTTCG	AATATCTACT	1260
TCCTTGATTT	TTACCGCCAC	CCCTTTATTAA	GCAAGAAGGA	AAACTCCTGC	TTCAAAACAAA	1320
CCACTGTAAA	GAACAAGCCA	CCCAATAGAT	ACGATAGAGA	TTTGTAAAAA	TGTCCTAA	1380
AGAATATCCA	ACACACTACT	CAAGAAAATA	ACAAAAAAATA	ATCTGT	TATTTTCTACT	1440
ACCTCCATT	ATTTATTTCA	CTAACAAATT	AATAGAGCCT	TCTACTCAA	TATCCTGTCA	1500
GAAAAGGATA	GAAAGCTACT	TTTTATAATA	CTTCAAGCCC	CACATGAGCA	GAAGCGTGAT	1560
AAACAAGCAG	AGAATACACC	TATATAAGCC	ATTAGTTGTT	GATAGAATT	TGTTTCTGAA	1620
ATACCTCTAT	ACAAACAAAT	GACAAACATA	AAATCTGCCA	AGCCGATAAA	CATAAGTTGA	1680
TTGCTTCTAG	GACTAACCAA	ATCATCATTT	ACTTATATTT	AAGAGTATCT	CTTTTATTTT	1740
AATGTATGTT	AGCACTGAAA	AGCAAGACAG	GCCAATAATA	TTTAAATGA	ACAGTAACGG	1800
GGTTAAGTCT	CTAAAAAAAT	TATCTACTGA	CACTACAAGA	AATACTATAC	ATATTATAGT	1860
CGAAACTATC	TTTTCTTAT	CCATAATTAT	TTACTCCTT	CCTAACAAAT	CCAGCTTATC	1920
AATCAAGAGC	GATTTTTAAC	ATAATGTAGC	AGCACCCGTT	GCAACTTTGA	CAAGTTAGT	1980
ATATCATTGT	TTTTAAAT	TTTCATCCA	AATCTTGAAT	TGTCATCGAA	ACATCTTGA	2040

246

TTGTTAAAAA	ATTTAAAAG	TAAGCATTAA	AAACATACTT	TCCTCTTTAT	ATTGTATTGA	2100
TACCAACTG	TTTGTAGACT	TTTCATCCTG	CTATCACATA	TCATTTGAC	AGGC GAAACA	2160
ATATTTAAAGA	AACTCCCTG	TAATTAAGC	TAGCAAATAC	AGGGGAGAAA	TTTATTTTTT	2220
AGAGAGTACT	ATCCGTATCC	TTTTTGGAAAG	ATTTGAAA	TATTTTCTA	ATTAAGTCAT	2280
CCATATAAGG	ACCAATATA	CCAACTACTA	AACCAATAAT	AAAACTTTA	AAATCCATAA	2340
TTACCACCAA	CATATTGCTG	CATAGGCTAC	ACCTCCAAGT	ATAGCTCCAC	CTGCAGCAC	2400
AGTTACACCT	ATTCCTATAG	CAAATGGTCC	CAATAGAAAT	GTCAAACCGT	TGTTGCACAC	2460
CCATCAATTG	CGCCATATGC	AACCCCTGCT	GCACAACTAA	TTTTCTTCC	CCAATCAATA	2520
TCTCCACCTT	CAACGCAAGC	AAGCATTCA	TTATCCATAA	CTGCAAATTG	TGACATCATT	2580
TTTGTATCCA	TATAGTGTAT	CACTTTTCAG	TTACGGAACA	AGTTTAATAT	AAAAATTATC	2640
AAAAAAACAT	AGGCAATAAA	GAGAAAATT	ATTTTATCAT	AGATTAGAAA	TAATATGACA	2700
AAACAAATTCA	ATGATGTTAA	TTCAATAGTC	TTTTGTTTTT	TATCGGAGAT	ACTTATGGAT	2760
AGATAAATAA	GATAGGTTG	AAAAGCGAAG	AGAATAATAA	AGAATATAGC	CTTCATAAAA	2820
TTTAGCTTTC	ATTTTATGA	TGTAGCGGTA	TAGGCTAAAT	ATCCACAAAC	CACTGCTCCT	2880
CCAATTCTC	CTATTGCAGC	GCCCCATGGT	CCTAGAAGTC	TCCCATATT	CACTCCACCC	2940
GCTGCACAAC	CTAAAGCAGC	AACTACAGCT	GCTCCTCCGG	AATTACCTCC	ATAAACCTCA	3000
CTCAGCATTG	TTTCATTAT	ATTACAATAA	GTATTCTAC	AACTCTCCTT	TTATTAAAAT	3060
CCACCCGTTG	CCCCCTGTTAC	TCCTGCCAA	AGATCCACAC	CAAATTAGC	TCCTATGTAT	3120
CCACATGCTC	CCATAAATGG	TGCTCCAACA	CCACTCGCAG	CACAAATAGC	TGTCCTAGC	3180
CCCCAGCCAC	CAAAGCAGC	ACCACCACT	TCTAAGACAT	TAGTTGCCA	ATTATTCTTG	3240
CCTCCTCAA	TACTAGATAA	CATAGTTATA	TCCATTTCAT	GAAATTGTC	CATAATTTT	3300
GTATCCATGA	CAAATACTCT	TTTTTATTTT	TAATTTTGT	TTGTTGTAA	CTTTGACAAG	3360
TTTAGTATAT	CATCGTTTT	AAAAATTTTT	CATCCAGATT	TTGAATAGTC	ATCGAAACGT	3420
CTTGAATTGC	AAAATTACA	TTAGACTTCC	TGCAAAACTA	GAATCCTAGT	TCATGATTGA	3480
TAATACCAAGC	ACTCAAATTC	ATTCGTAATC	CGAAGCGTTT	ACGATGACTT	CGATAGGTTG	3540
TTGAAAACAT	TTTAAACGTT	TTTACTTTGG	CAAAGATGTT	CTCAACCTTG	CTTCTCTCCT	3600
TAGATAGCGC	ATGGTTACAG	GCTTTATCTT	CAACTGTTAG	CGGTTTGAGT	TTGCTGGATT	3660
TACGTGAAGT	TTGTGCTTGA	GGATATATCT	TCATGAGCCC	TTGATAACCA	CTGTCAGCCA	3720
AGATTTTACC	AGCTTGTCCG	ATATTCTGC	GAETCATTT	GAACAACTTC	ATATCATGAC	3780
AATAGTTCAC	AGTGATATCC	AAAGAAACAA	TTCTCCCTTG	ACTTGTGACA	ATCGCTTGAG	3840

TCTTCATAGC GTGAAATTC TTTTACAG AATCATTGCG TAATTCTTT TTTAGGGCGA	3900
TTGATTTTA CTTCCGTCGC ATCAATCATT ACCGTGTCCCT CAGAACTGAG AGGAGTTCTT	3960
GAAATCGTA CACCACTTG ACAAGAGTT ACTTCAACCC ATTGGCTCG ACGGAGTAAG	4020
TTGCTTTCGT GAACACAAA ATCAGCCGCA ATTTCTTCAT AAGTGCAGTA TTCTCGCACA	4080
TATTGAAGAG TGGCCATAAG AAGGTCTTCT AGGCTTAATT TAGGTTTCG TCCACCTTTT	4140
CGCTGTTAA GTTGATAAGC TGTTTTAAT ACAGCTAGCA TCTCTTCAAA AGTCGTGCC	4200
TGAACACCAA CAAGACGCTT AAATCGTGCA TCAGTTAGTT GTTACTTGC TTCATAATT	4260
ATAGAACTAT AGTAAAATGA AATAAGAACAA GGATAAATCG ATCAGGACAG TCAAATCGAT	4320
TTCTAACAAAT GTTTAGAAG TAGAGGCCGA CTATTCTAGT TTCAATCTAC TATACTATAC	4380
CATATTTGT TTGCGAGGGAA ATCTATTATA AAAGGTAAG TATTGCAAAA ACACCTACCC	4440
TTTTCTTTA TACTTCATTA AGCTCTACTT TTATATAAC TTCAAGCCCC ACATGAGCAG	4500
AAGCATGATG ATTAAGCAGA GAACAGCGCC AATATAAGCG ATTATTTGTT GGTAGGATT	4560
TCCTGCTGTG ATACCTCTAT ACAAAACAAAT AATAGACATA AAACCTGTCA AGCCGATGAA	4620
CATAAGTTGA TTGGTTCTAG GACTAACCAA ATCATCATCT TCAAACCTCTC TTATCCTCAT	4680
TTCCCTAGTG AGATAAACAG TAACCAAAAT AGAAGCCAAG TTAATAACTA CTAAAAGAAA	4740
TTGGAAAAT ACGGAAAAT TTAAAAACTG ACGAGATAGA AATAGATAAG TAGAAACAAAG	4800
CAAGGGCAAC TGACCTAAGA ACAATCTCGC AAGGAAGATG TTCCGTTTT TAGCAAGAAA	4860
AGTTTCATT TCTTTCTCC TTTCTTTTA TTGATAGCAA AATAGATCAT AACTGCAATC	4920
ACATAGGCTA TGGTATAAAA TAGCTGATAC CAAGCACTCT CCCTAACCGG ATATAGAAAAG	4980
ATGGACATGA TTAGATACAG AACGAAAATA ATCACTTATT TTTCTTCAT AAGATTTCT	5040
CCTAAATGTG CGATTTATCT TAGTTGAGCA AGAACATTAA CACTGCTAGT ATAGCACTTA	5100
TTTGACCTT GGATCACTCA AATCATAAAT GGTCACTAAA ACCTCTTGAA TTGTAAGAAAAT	5160
TAAAAAAAGCA AGCATGAAAA ACATACTTTC CTCTTTATAT TGTATTGATA CCAACTTGT	5220
TGTAGACTTT TCATCCTGCT ATCACATATC ATTTTGACAG GCGAAACAAT ATTAAAGAAA	5280
CTCCCCGTGA AATTAAGCTA GCAAATACAG GGGAGAAATT TATTTTTAG AGAGTACTAT	5340
CCGTATCCTT TTTGGAAGAT TTTGAAAATA TTTTCTAAT TAAGTCATCC ATATAAGGAC	5400
CAAATATACC AACTACTAAA CCAATAATAA AACTTTAAA ATCCATAATT ACCACCAACA	5460
TGTTGCTGCA TAGGCTACAC CTCCAAGTAT AGCTCCACCC GCAGCACCAG TTGCTGCACC	5520
TTGCCATGTT CCTGTTTAA TGCTAGTTG AAGACCTCTT GCTGCTCCTC CTCCAACACC	5580

248

TGCTTTGGCA	AAATCTCCCC	AATTGCATCC	GGCACCTTCA	ACGCAAGCAA	GCATTTCA	5640
ATCCATAACA	AAAAATTGTG	ACATCATT	TGTATCCATG	ACAAATACTC	CTTTTTAA	5700
AAACTAAAAT	AAATCAGAAT	AGAATCCTCA	TAATTTTACT	ATAAGTCTTA	CCAACTTAGT	5760
CCCAATTAT	CACCAACCAT	ACCTCCTAAG	CATGTTAATC	CACCCCAAT	TGCACCAATG	5820
TGTGCTCCAA	CAAATGCACC	AGCAAGTCCA	GCTACTCCTA	AACTGGCCAA	ACCTGCTCCA	5880
GTTCCACCAAG	TTATAATTCC	CGTAGTGACT	CCTGTAATCA	GTGCAATTG	ACAATCAGTG	5940
GAGCTATACC	CCCCTTCAAC	TTTCGCAAGC	ATTTCA	CCATAACCTC	TAACGTGAC	6000
AACATTTTG	TATTGATGAT	GAATACCTCC	TTTTTATTTT	CAATTGTTA	CCAAAGTCTT	6060
AAATTCAATA	AAACAAATAGA	TTTTTTATAG	TATCTTTTG	ATTTCTTAA	AAAAGTATAT	6120
ACGTCTACTA	TCTTCTTAA	GGTAGCAGTA	CCTATTTTTT	AGTCTAAGAT	TTCAATAATC	6180
TTGAGTATCT	AAATATCTT	AATTGCGTTA	TTCTCCTTGC	AATAAAAGT	TTTACTATAC	6240
TATTTATTAA	CTTGAGAAA	GCAAAAATA	TTAGTAAATA	ATAGTTATA	GTAAAGTTT	6300
TTATTCCTAC	CAATCCATCA	ACTAAGTAAA	GCATCAACGA	TTACATAAAC	GATTGATAAT	6360
ATAATTAAAA	TTTGCTAAC	TATCTTATTC	TCATCATTCT	TAGATAACTT	TGATATTG	6420
TAAGTAAGTA	AATAAGACAG	AAATTAATA	GCGATAATAA	TACTATATTT	AAGAATCATA	6480
ATCTTACAAA	GAGGACATAA	TTCCTGAACC	TACACAAATA	AGTGTGCTG	CTCCCCAGT	6540
TATCGGACCA	GTGCGAGCAG	CTAATAGTAC	TGCTCCAATA	CAACCACCGA	TTGCAGATCC	6600
TAAATTGCGCT	CTTCCTCCAC	TAACTATTTC	GAGTTCTTCA	TTATCCATAA	CAGAAAATTG	6660
TTCCCATCATT	TTTGATTCA	TGACAAATAC	TCCTTTTTC	TTTTTTATT	TTTGTCTTGT	6720
TGTAACTTG	ATAAGTTAG	TATATCATCG	TTTTTTAAA	TTTTTCATCC	AGATCTTGAA	6780
TTGTCATCGA	AACGTCTTGA	ATTAGCTTT	TTATTCAG	CCACCTCTAA	ATGTTAAAA	6840
AAAATAATT	CTAACACTT	TTTACCAATT	CAGGAAGTTT	TAATGACTAT	TCAAGATTTC	6900
ATAAAATATG	AACTTAGTTT	TATGACATAA	TAGACCTATC	CACTATATGA	AAGGAATTGC	6960
CAATGACTTC	TTATAAACGT	ACATTTGTC	CTCAAATAGA	TGGGAGAGAC	TGTGGTGTG	7020
CTGCTTAGC	CTCGATTGCT	AAATTCTATG	GTTCAGATT	TTCTCTAGCT	CACTTGAGAG	7080
AACTTGCAAA	GACCAATAAA	GAAGGGACGA	CTGCTCTGG	CATTGTAAA	GCCGCTGATG	7140
AAATGGGCTT	TGAAACAAGA	CCTGTTCAAG	CAGATAAAAC	GCTCTTGAC	ATGAGTGTG	7200
TCCCCCTATCC	ATTTATCGTT	CACGTTAACCA	AAAAGGAAA	ACTCCAACAT	TACTATGTTG	7260
TCTATCAAAC	AAAGAAAGAC	TATCTGATTA	TTGGTGATCC	TGACCCCTCT	GTAAAAATCA	7320
CTAAAATGTC	AAAAGAACGC	TTTTCTATG	AATGGACTGG	AGTAGCTATT	TTTCTAGCTA	7380

249

CCAAACCCAG	CTATCAACCC	CATAAAGATA	AAAAGAATGG	TCTACTAAGC	AAGCTTCCTT	7440
CCTCTGATTT	TCAAACAAAA	ATCTCTCATT	GCTTACATTG	TTCTCTCAAG	CTTATTGGTC	7500
ACTATTATCA	ATATAGGTGG	TTCTTACTAT	CTCCAAGGAA	TCTTGGATGA	ATACATTCCA	7560
AATCAGATGA	AATCAACTTT	AGGAATCATC	TCAGTTGGTC	TGGTTATCAC	CTATATCCTC	7620
CAACAAAGTC	TGAGCTTCTC	CAGAGATTAT	CTCCTAACCG	TTCTGAGTC	GAGATTAAGT	7680
ATTGATGTGA	TTTTATCCTA	TATTCGCCAT	ATTTTGAAAC	TTCCCATGTC	TTTCTTTGCC	7740
ACACGTCGTA	CAGGAGAAAT	CATTCACGA	TTCACAGATG	CTAACTCTAT	TATAGATGCC	7800
TTGGCTTCTA	CCATTCTTC	TCTTTTCTG	GATGTTCTA	TTCTGATTCT	TGTAGGAGGC	7860
GTCTTACTGG	CACAAAACCC	TAATCTCTTC	CTTCTTTCTC	TTATTTCCAT	TCCTATATAC	7920
ATGTTCATCA	TCTTTTCTTT	TATGAAACCT	TTCGAAAAAA	TGAACCATGA	TGTCATGCAA	7980
AGTAATTCTA	TGGTTAGCTC	TGCCATTATC	GAAGATATCA	ACGGGATTGA	AACTATAAAG	8040
TCGCTCACGA	GTGAAGAAAA	TCGCTATCAA	AAATAGACA	GCGAATTGT	AGATTATTTG	8100
GAAAATCCT	TTAACGCTCG	AAATATTCT	ATTTACAAA	CGAGTTAAA	GCAGGGAACA	8160
AAATTAGTTC	TGAATATCCT	TATCCTATGG	TTGGCGCTC	AAATTAGTCAT	GTCAAGTAAA	8220
ATTTCTATCG	GTCAGCTGAT	TACCTTTAAC	ACACTTTTT	CTTACTTTAC	AACTCCTATG	8280
GAAAATATTA	TCAACCTCCA	AACCAAACTC	CAATCTCGA	AGGTCGCTAA	TAACCGTTG	8340
AACGAAGTCT	ATCTAGTCGA	ATCTGAATT	CAAGTTCAG	AAAACCCCTCT	TCATTCACAT	8400
TTTTGATGG	GCGATATTGA	ATTTGATGAC	CTTCTTATA	AGTATGGTT	TGGATGAGAT	8460
ACCTTAAACAG	ATATTAATCT	CACGATTAAA	CAAGGAGATA	AGGTTAGCCT	AGTTGGAGTT	8520
AGTGGTTCTG	GTAAAACAAC	TTTAGCCAAA	ATGATTGTCA	ATTTCTTTGA	ACCCTACAAA	8580
GGGCATATTT	CCATCAATCA	TCAGGATATT	AAAACATTG	ATAAAAAAAGT	CTTGCGCCGT	8640
CATATTAATT	ACCTACCCCA	ACAAGCCTAT	ATCTTTAATG	GCTCTATTTT	GGAAAACCTTA	8700
ACCTTGGCG	GTAATCATAT	GATTAGTCAA	GAAGATATTC	AAAAGCTTG	TGAAGTAGCT	8760
GAAATCCGTC	AAGACATTGA	AAGAATGCCT	ATGGGCTATC	AAACTCAGCT	CTCTGATGGA	8820
GCTGGTCTAT	CAGGAGGACA	GAAGCAACGA	ATCGCTCTCG	CTCGTGCCTCT	TTTAACTAAA	8880
TCTCCTGTTT	TAATACTAGA	TGAAGCTACT	AGCGGTCTTG	ATGTCTTGAC	TGAGAAAAAG	8940
GTTATAGATA	ATCTTATGTC	TCTAACTGAT	AAAACCATTC	TCTTTGTAGC	CCATCGTCTC	9000
AGTATAGCCG	AACGAACCAA	CCGTGTCATT	GTTCTTGACC	AGGGGAAAT	CATTGAAGTT	9060
GGTA						9064

250

(2) INFORMATION FOR SEQ ID NO: 18:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 7780 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 18:

CTCCATTTTT TTGATTCAT AAATAAACAA CCTCTCTGTT AATTTGTAT AATTATAACG	60
ATATCCAAGT TACTTGTCAA GTGTTTTTA AATTTTATC TCAAAATAT TTTTCGTTT	120
AAAAAAAGGA GCCATCAGTT GATTCAAGC TCCCTTTAT ACAGAATTAA ACTATTTAT	180
AGTTCGACAA TCTTACCTGT TTCAAAGTAG ACAACCCATT CACAGATATT TTTAGCATAG	240
TCACCGATAC GCTCCAAGTA GGAAAATAACT TGGAAAATAAT CACGACCCGT AACAAATGGCT	300
TCTGGATTTT TCTTAATCTC TTCAGTCGCA AGGTACCGGA TAGTTCAAA ATAGTGGTTA	360
ATTTGCTCAT CCATGGAGGC CACCCGGTAT GCGTCGTCAA CAGAACATT AAGATAAAGA	420
TCAAGTGCTG CTTCCACAAC GCTTTAACT TCACGTCCA TTTTTTTAAT TTCTTCCTCT	480
ACAGCTGGAA TGCCTCTTC CCCCTTCATA CGGATGGTTG CCTGGGCAAT GGCTACAGCG	540
TGATCCCCCA TACGCTCCAC ATCTGATACA GCCTTAAGGA CAGTCAAGAC TGACGCAA	600
TCTTGAGAGA CTGGTTGTTG GAGTGCAGTC ATTTCAGATG ATTTCAGTTTC CAGTTTCACT	660
TCTGATTCAT TTACTTCTGC ATCATCTTCG ATGACCTCTT TTGCCAGGTC ACGGTCATGC	720
GTGACAAAAG CACGTACCGT ACGATTGATT TGTGAGAGCA CTTCTTGTC CATAAGCGTAG	780
AACTGGTTAT GTAATTTCTC TAAATCTTCT TCAAATTGAG ATCGTAACAT CTTTCATCTC	840
CTTATCCAAA TTTTCCTGTA ATATAGTCTT CCGTTCCCTT GTGTTGGGGA TCAAGGAACA	900
TCTGCTTGGT ATCATTAAAT TCAATCAAAT CTCCATCTAG GAAAATCCT GTCTTATCAG	960
AGATACGTGA AGCTTGCTGC ATGGAACGGG TTACCAAGAG CATGGTGTAC TTGTCTTTA	1020
GACCATACAA GGTTCCCTCA ATTTTACCAAG CTGAAATCGG ATCCAAAGCC GAAGTTGGCT	1080
CATCCAAGAG GATGATTTA GGACTAGTTG CCAAGACACG GGCCACGCAG ACACGCTGCT	1140
GTTGACCACC TGACAATCCA ATAGCTGAAT CATATAGACG ATCCCTGACC TCATCCCAGA	1200
TAGAGGCACC TTGCAAGGCT TTTTCTACGG CTTCATCCAG AACCTGCTTA TCCTTAATTG	1260
CATTGATACG AAGCCCGTAG ACAACATTCT CATAGATAGT CATAGGGAAA GGATTAGGTT	1320
GTTGGAAAAC CATTCCGATT TCCTTACGTA ATTCAACCGT ATCTGTACGC GGACTGTAGA	1380
TGTTGTGACC ATTGTACACC ACGGATCCAG TTGTGGTCAC CTCTGGATTG AGATCTCCCA	1440

TGCGGTTGAG AGACTTGAGG AGGGTTGACT TCCCCTGATCC AGATGGACCA ATCAAGGCTG	1500
TAATTCCTT AGGTTGGAAA GATAGGGAAA CACTATTCAA AGCCTTCTTT TTATTATAAT	1560
AAACGGACAG GTCTGATACC TGTAAAATCG CACTCTGTAT ACAGGTTTCCT TTCTAACCAA	1620
AGTGACCAGA TACATAGTCA TTGGTGGACT GTAGCTTGGC ATTTTGAAA ATAGTTGCAG	1680
TCTTGTCTATA CTCAATCAA TCACCCAAGT AAAAGAAGCC TGATAGTCA CTTGCACGAG	1740
CAGCCTGCTG CATATTATGC GTTACAATGA TGATGGTAAA GTTTTCTTG AGCTAAACAA	1800
TGGTCTCTTC TAGTTGCATG GTCGCAATCG GATCCAAGGC TGAGGCTGGC TCATCCATTA	1860
AGAGGATATC TGCGTTAACCA GAGATGGCAC GAGCGATACA GAGACGTTGT TGCTGACCAC	1920
CTGATAAGGT CAAGGCTGAC TTGTTGGAGAT CGTCTTTAAC CTGATCCAG AGGGCAGCCT	1980
GACGAAGGGA GGTTTCTACG ATTCATCTA GGACTTGCTT ATCCTTAAC CCAGCACGTT	2040
CATGCGCAAA GGTAATATTA CGGTTAAATTG ACTTAGCAAA TGGATTGGGA CGTTGAAAAAA	2100
CCATTCCAAT GTGTTTACGC ATTCATAAA CGTTGATTTC TGGACGGTTG ACATCAATT	2160
CACGATAGAG AATCTGCCA GTTACTTTAG CAATATCAAT AGTATCATTC ATGCGATTGA	2220
GACTGCGTAA GTAGGTAGAT TTCCCCGATC CCGACGGGCC AATCAAAGCT GTAAATTAT	2280
TTCTTCTAAA TTGCATATCA ATCCCCTAA TGGATTCTT TTTACCATAG TAAACATGGA	2340
CATCCTTAGT AGAAAGGGCT ACTTTTCTT CAGGAAAGGT AAGGATATGC TTCTCATCCC	2400
AGTTATATGT TGACATGGCT TCTCTTTAG GCAGCGGTTA ATTTCTTGTG TAGATACCTT	2460
CCGAACCTAC GAGCTCCAAA GTTAAAATC AGGATAAAGA TCAGGAGCAC AGCGGCAGAA	2520
CCTGCTGATA CAATGGTCC ATCTGGAATA GTGCCCTCAC TATTGACTTT CCAGATATGG	2580
ACAGCCAAGG TTCTGCTTG ACGGAAGATA GAGATGGGCC TAGTCACACT GAGGATATT	2640
CAGTTAGACC AGTCAAGAGC TGGCGCCGAT TGCCCTGCTG TATAGATCAG AGCTGCAGCT	2700
TCGCCAAAGA TACGACCAGA TGCCAAGACG ACACCCGTTA CAATACCTGG AAGCGCTTCC	2760
GGAATAACAA CATGAACCAC TGTCTCCAG CGAGAAATCC CAAGAGCCAG ACCAGCCTCA	2820
CGTTGGGTAT GGTGAACCTG TTTCAAACCA TCCCTCTACAT TACCGCGTCAT CTGAGGCAAG	2880
TTAAAGACTG TCAAGGCCAA GGCACCTGAA ATGATTGAAA ATCCATACTC AACTGGACT	2940
ACAAAGATCA AGTAACCAAA GAGACCCACC ACCACTGATG GTAAAGAGGA CAAAATTCA	3000
ATACAAGTCC GCACAAAGTT GGTAACAGGA CCTTTTTAG CATATTCAAG CAACTAAATC	3060
CCAGCTCCCA TAGAAAGAGG TACAGAAATA ATCAAGGTAA TGACCAATAG GAAAAGGAA	3120
TTGTAAAGCT GAATGCCAAT CCCACCCACCT GCTTGAAAAG CAGAAGACCT TCCAGTCAG	3180

252	
AAAGACCAAG AGATATGGGG CAAGCCCCGA ACCAAGATAT AGAGAATCAA GGAAGCCAAG	3240
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AACTAAGCTC ATCAAGAGCA GTACCAAGGC CAGTGACCAAG AGAACATTAT TATTTACAGT	3420
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ACCAAAGGCA CGGCCATCC CAAAGACCAC TGCACTGAAA ATACCAAGAAC GGGCCGCCTT	3600
CAAGATCACA CGCCAGATAG TCTGCCAGCG AGTGGCTCCC ATAGCAGAACAC TGGCTTCACG	3660
ATAATAACGA GGAACCGCAC GCAAGCTATC CGTTGTCATA AAGGTTACGG TCGGCAAAAT	3720
CATGACAAAG AGGACGGAAA TCCCTGACAA AATCCCCAAA CCAGTCCCAC CAAAGACACT	3780
GCGAACAAAG GGAACCGACGA CTTGCAAGCC AATAAATCCG TACACTACTG AAGGAATCCC	3840
AACCAGGAGT TCAATAGCTG GTTGCAAAAT CTTCGCCCTT TTTGGTGATA CTTCGGTCT	3900
AAAAACTGCT GCACCAATAG CAAAGGGTGT TGCGATAAGG GCTGAGAGAA TGGTAACGAT	3960
AAAGGAACCC AAAATCATAG GAAGGGCACC AAATTCTTTA CTAGAAGGAT TCCAAGTTCC	4020
TCCCCAAAGA AAGTCAAAGA TATTCAACACC ATTGACAAAG AAGGTCGACA AGCCTTTTG	4080
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CAAACCTTTT CCTAATTCTC CCAGACGAGA ATTCTTGAT GGAAGCAACA TTTCTTAGC	4200
TAATTCTTCT TGATTCTTCA TTGTCCTCCCT TCCAACACTG TCACAGTTCC GGCAGCATCT	4260
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GTCTCATCCG AGAGAACAAA ATTGAGAAAT TCTGCAGCCA ACTCATTGGG CTGCCCAAT	4380
GTATACATAT GCTCATAAGA CCACAAGGGC CAATTATTGC TACTTATATT TTCTGGACTT	4440
AAAGTCATAGC CATTCAACTT CATGCTTTTG ACCGAATCAT CTATATAGGT AAGAGATAAA	4500
TAAGAGATAG CTCCTGGACT TTTTGATACG ATTGATTTTA CCGCTCCATT TGAATCCTGC	4560
TCCTGACTTT GCATGGCAGA CTGACCTTCC ATAATGACAG TATCAAAGGT ACCACGAGAG	4620
CCAGAGCCGG CTGCCGATT GATAACAGAG ATGGTAAGT CCTTACCAACC AACCTCTTTC	4680
CAATTGGTTA CCTCACCTAT GAAGATTGAGA CGAAGTTGCT CTGTCGTTAG GTTATCAACA	4740
TCAACCTCCT TATTGACAAT CAGAGCCAAG CCAGCTACCG CGACCTTGTG CTCAACAAGA	4800
GCAGAACAT CAATTCCGTC TTTTCCTCA GCAAATACAT CTGAGTTCC TATATCAACT	4860
GCCCCAGACT GAACCTGGGA CAAGCCTGTA CCAGAACCTC CCCCTGGAC ATTGACCGTT	4920
TTTCCAACAT GGATCGTGCC AAATTCTATCT CCCGCTACTT CAACCAAGGG TTGCAAGGCA	4980

GTTGAGCCAA CAGCCGTTAT GGATTCTCCA CGATCAATCC AGCTAGCACA GCCTACTAAA	5040
CAAGCCGTCA GCCAAAAGC GATAAGAGAC AGAGCAAGCT TTTTTCTTTT TTTCACTGTT	5100
TTTCTCCTCG AAAATAATTA TGAATACTGT GAATTTTTA AGTAGTTCTT TATGAGTTGA	5160
CGCATGAATT CTTACCAAAT TTCTGCGCAA TTGATTATTT ATATAATATA GGCTATATTA	5220
CTCTTCTCTA ACCTCCTTT TTTCATATGTG GATAAAATCT CTTGTCTATC CCTTCCCCCA	5280
TTGTCACCCA TTATAGTCAT TTCTGTCTC TTTTTCCCT TTTTAATGCA AGGGAAATTA	5340
CTCTCCTTAG ATGATAATCC AAAAGCTAGA AAGGTATCTC AAACCTCTCT ACTCTCCCAG	5400
ACTAGTTAC AACTAAAAGG AAAAGATTCT ATTTTATGAG AAATCTAGTT TACAAGCGGT	5460
AAGAACGCTA ATAACAAAC TTCTGTACT CTTTGAAAAT CTCTTCAAAC CAGTGTGTTG	5520
AGCTATCTAT GGCTAGCTTC CTAGTTGCT CTTTGATTT CATTGAGTAG TAAAACATACA	5580
TGTAATGGCA ATCAAGATAT CAAGAATCAT CCTACTAAA AAATCCATAC TTTCACTATA	5640
ACATAGAATA AGATATTGTA CTAGCATTCTT CATTGAAATC TGAGGCCTTT TGAAAATAA	5700
TTTTCAAAA CATTCCAGT AACCTTTGCA AAGCCCAAGC CATTGCCTTT ACCAAAAACT	5760
TGGTACCAAC CATTGGCAG ACTTTCTGCC AGCTGAACGG TTTCTCCAGC CGCATACTTG	5820
ACAAACGCTT CTTGGCAAT TTCAACCGAC TGTTGACCT GACTCGGTTT CAAGGCTAAA	5880
CCAAGAGCGA AACTGGGCTC AAAGCGTTTC TTCTTAAAG TACCCAGATG CAGTCCATTG	5940
CGAGCAATCT TGAGCTTCCA TAAATCTGGC AAAAGTTCTG CCAAGAGATA AAGCTGGCT	6000
CCAAAAATCT GCAAGATACC CGGTAGATTG ACCTTCAAAT GTTTTGGGC AAATTCTGC	6060
CACAAGGCAA CTTGTTCACG GCTGAGGTTA CTCTTACTTG CCTTAAATTT AGGAGCTGGA	6120
TTGTTACCT TAAACTGTAG ATGGGCAACA AACTGACCT CTCCCTTAAA CTGATGAGGA	6180
TACATCCGAG CGCTTCTCG CAGGTCAATA CCAGCTACCA TTCCATTGAT ATGCTCTACT	6240
GGCAACAAGT CAAAATCATA CTCTTCCAGC AACCAATTGA CAATCTCTC GTTTCCCTCG	6300
GGTGCCCAGG TACAGGTGCA ATAAACCAGA TGACCACCTT CAGCTAACAT GGTCACTGCA	6360
TCCTCCAGAA TTCTCTTTG CAAGCTAGCA CATTGACTCG GATAATCTAA GCTCCAATAG	6420
TCCATAGCAT CAGGTGCTT ACGAAACATT CCTTCACCGAG CGCAAGGGGC ATCAAGAACG	6480
ATTAAGTCAA AATAGCCTTT AAAGACCTTG ACCAAGCGGT CGGCAGATTG ATTGGTCACC	6540
ACGACATTG TCGCTCCAAA ACGCTCCATG TTTTCAACCA AAATCTTAGC CGGTTTGCTT	6600
GAAATTCAT TGGAAACAG TAGCCCCCTCC CCTGCTAGAT AGGCTGCCAG TTGAGTTGAT	6660
TTGCCCTCCCG GTGCAGCAGC CAAGTCCAAG ACCTTCATAC CAGGACTGGG TTGGGCTACT	6720

254

TGAGCCACCA	TTTGAGCAGC	AGGTTCTTGC	GAATAAACTA	AACCTGTAGC	ATGCTCAGGC	6780
GATTTCCCTG	AAACCTTCCC	ATAGTGGCCC	CAAGGGTTT	GAGTAATGCC	ATCAGAAAAG	6840
GAAAGTTGCT	CTTCCTTTAA	GGGATTGACC	CGAAAGGCCG	AAACCGCTTC	CTCCTCAAAA	6900
GAGGCAAGAA	AATCTCTTGC	CTCATCTCCT	AGTATCTCTT	TATATTTTC	AACAAATCCT	6960
TCTGGAAATT	GCATTTAAGT	TCTTTCCCTT	TCGTAATAT	AGGACTGAAT	TTCCCTCCTGC	7020
ATCTCAAGAG	GCACCATCAT	GACCGGCTGT	CTGGTTGAA	AATCAGGAGC	TTCACCAAAA	7080
AGGGTCACAA	CCCGATAGCC	CAGACTTTCC	CCTAAAATAC	TAGCTGCGGC	ATAATCCCAT	7140
GGTTGCAGAT	AAGTGGAGATA	GGTCAACAAA	CGCCCTGACA	AAATCTGGC	AAAACATAATG	7200
GCCGCACTTC	CATAGACACG	AACACCAAGA	ACCGCTCGGC	TCAAATCAGC	CAGCCCCCAT	7260
TCATTGGTTT	CCAGCATAAC	ACTATTCCCT	GCAATGAGAA	AATCTCCAAG	TGGTTTAGTT	7320
TTAAAAGGAG	CTAGGGACCT	ATCATTAGA	CAAACCTGGAA	ATTCCCCACC	ACCGTGGTAA	7380
CAATCCCCTT	TGACCACATC	ATAAATCAGA	CCAAACTGTC	CCTGACCATT	TTCAAAATAA	7440
GCCATCATAA	CAGCAAAATC	TTCCTGCTGG	GCTACAAAAT	TATTGGTACC	ATCAATGGGA	7500
TCAATGACCC	AAACCTTGCC	CTCTTGAAACC	GAGGCTCGCA	GACAACCTTC	TTCAAGCACAA	7560
ATCTTATCCT	CAGGATAACG	GGACAAAATC	TCACCAACCA	AGAGTTCTG	AACTTCTTIG	7620
TCCAGCTGG	TCACCAAATC	TGTTGGAGAG	GACTTGGTTT	CAACACGCAA	GTCTTCCCTGC	7680
ATATGGTCAA	GAATGTACTG	ACCTGCTTTC	TTAACAAAGCT	CTTTAGCAA	TTCAAATTTA	7740
CTTTCCAAGA	GAAATCTTTC	CTTCCCCTTT	TTCTTTGGGG			7780

(2) INFORMATION FOR SEQ ID NO: 19:

- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 4820 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 19:

GTAATGATAT	AGGAACACCA	GGTGACCTGA	TGGGACGTCG	TAAGCCTATG	AACTACTAGC	60
TGCTAAAGGC	TTTAAAGATG	GTATGGTACC	ATATATCTCA	AACCAATACG	AAGAAGAACG	120
CAAACAAAAG	GGCAAGACAA	TCAATCTCTA	CGGTAAAACA	AGAGGTTTGG	TTACAGATGA	180
CTTGGTTTTG	GAAAAGGTAT	TTAATAACCA	ATATCATACT	TGGAGTGAGT	TTAAGAAAGC	240
TATGTATCAA	GAACGACAAG	ATCACTTGA	TAGATTGAAC	AAAGTTACTT	TTAATGATAC	300
AACACAGCCT	TGGCAACAT	TTGCCAAGAA	AACTACAAGC	AGTGTAGATG	AATTACAGAA	360

255

ATTAATGGAC	GTTGCTGTT	GTAAGGATGC	AGAACACAAT	TAATACCATT	GGAATAACTA	420
CAATCCAGAC	ATAGATAGTG	AACTCCACAA	GCTCAAGAGA	GCAATCTTA	AAGCCTATCT	480
TGACCAAACA	AATGATTTA	GAAGTTCAAT	TTTGAGAAT	AAAAAATAGT	GTCTACTATT	540
AGGAATAAAA	GTTTAAAAG	GTGATGAAGA	ACAAACCAAG	ATTCAAGCAG	GAATTCCCTAC	600
TGATAATGAA	GTAAGTTATG	ATCTTATTTA	TCAGCAGGAA	ACTCTTCCTG	CAACAGGTC	660
ATCAACTTCT	GAGCTTACAG	CTTTAGGCCT	ATTAGCTGTT	GGTAGTTTAG	TTCTTTGGT	720
TCATAATATG	ACGGAACAG	TTTTTGCTC	CCTCTGAAA	GTCATCATTT	GATGGCTTT	780
TTCTATATAG	GGTAAAAGAT	AGGGTAAAAG	GCTATCATCG	GACAAAATAA	AGAAGGCATG	840
ATATAATATA	AAGTAGATTT	CTATGTCATA	AAACAAGAAC	TGTTTGGACA	TCATTCAATT	900
GAAAACCTCT	TATGTTCAA	CAATAGTAAA	ATAAAATAGG	GGATCTAAAT	CCTTGCTATG	960
AAAGGAAAAA	ACTCAATGGC	TAATTCAA	TGGTTTCCTG	GTCACATGTC	TAAAGCTCGT	1020
CGACAGGTGC	AGGAGAATT	AAAATTGTT	GATTTGTGA	CGATTTAGT	AGATGCACCG	1080
TTGCCCTCTAT	CTAGTCAAA	TCCTATGTTG	ACCAAGATTG	TTGGTGTAA	ACCAAAACTC	1140
TTGATTTAA	ACAAGGCGA	CTTGGCTGAT	CCAGCAATGA	CCAAGGAATG	GCGTCAGTAT	1200
TTTGAATCAC	AAGGAATCCA	GACGCTAGCT	ATCAACTCCA	AAGAGCAAGT	GACTGTAAAA	1260
GTTGTAACAG	ATGCCGCCAA	GAAGCTCATG	GCTGATAAGA	TTGCTCGCCA	AAAAGAACGT	1320
GGGATTCAAGA	TTGAAACCTT	GCCTACTATG	ATTATCGGGA	TTCCAAACCGC	TGGTAAATCA	1380
ACTCTGATGA	ACCGTTGGC	TGGTAAAAG	ATTGCTGTTG	TTGGAAACAA	GCCAGGGGTC	1440
ACAAAAGTC	ACAATGGCT	AAAAACCAAT	AAAGACCTGG	AAATCTTGA	TACACCGGGG	1500
ATTCTCTGGC	CTAAGTTGA	GGATGAAACT	GTTGCACTTA	AGTGGCATT	GACTGGAGCT	1560
ATCAAAGACC	AGTTGCTTCC	TATGGATGAG	GTTACCATT	TTGGTATCAA	TTATTTCAA	1620
GAACATTATC	CAGAAAAGCT	GGCTGAACGC	TTCAAACAAA	TGAAAATTGA	AGAAGAACCG	1680
CCTGTGATTA	TTATGGATAT	GACCCGCGCC	CTCGGTTCC	GTGATGACTA	TGACCGTTT	1740
TACAGTCTCT	TCGTGAAGGA	AGTCCGTGAT	GGCAAACCTCG	GTAACTATAC	CTTAGATACA	1800
TTGGAAGACC	TCGATGGCAA	CGATTAAGA	AATCAAAGAA	TTCCCTGTGA	CAGTCAGGAA	1860
GTTAGAAAGC	CCTATTTTTT	TAGAGCTTGA	AAAGGATAAT	CGCTCAGGAG	TTCAAAAGGA	1920
AATCAGCAAG	CGTAAAAGAG	CCATTCAAGC	TGAATTAGAT	AAAAATTGCG	GCTTGGAAATC	1980
CATGCTTCT	TATGAAAAG	AACTTTATAA	GCAAGGATTG	ACCTTAATTG	CAGGTATTGA	2040
TGAGGTTGGT	CGTGGTCCTC	TTGCTGGTCC	TCTAGTCGCT	GGGGCCGTTA	TTTTATCTAA	2100

256

AAATTGTAAG	ATTAAGGTC	TCAACGACAG	CAAGAAAATT	CCTAAAAAGA	AACATCTGGA	2160
GATTTTCCAA	GCCGTTCAAG	ACCAAGCCTT	GTCGATTGGA	ATTGGTATCA	TAGATAATCA	2220
GGTCATCGAC	CAACTCAACA	TCTATGAAGC	ACCCAACTA	GCCATGCAAG	AAGCAATCTC	2280
CCAGCTCAGC	CCTCAACCAG	AGCACCTTTT	GATTGATGCC	ATGAAACTGG	ACTTGCCCAT	2340
TTCACAAACC	TCCATTATCA	AAGGAGATGC	CAACTCCCTC	TCTATCGCAG	CAGCATCTAT	2400
AGTAGCCAAG	GTAACACGTG	ATGAATTGCT	GAAAGAACAT	GATCAGCAGT	TCCCTGGCTA	2460
TGATTTCGCT	ACTAATGCAG	GATATGGCAC	AGCTAAACAT	CTGGAAGGCC	TCACAAAAC	2520
AGGAGTTACC	CCAAATCACC	GAACCAAGCTT	TGAACCCGTT	AAATCACTGG	TTTTAGGTAA	2580
AAAAGAAAGT	TAATTGAAAG	GAAATAACAT	GGAGGAACAG	TCGGAAATAG	TCCGTTCTAA	2640
GAAAGAATTG	GCCTTTGCAT	CCAGCACTAT	ACTATCCAA	GTTGGTCAG	GAATCATTGT	2700
CGGCCTCATH	GTTGGAATT	TCGGCGGATC	CTTTCGTTTC	TTAATTGAAA	AGGGCTTCCA	2760
CCTGATACAA	GGAGTTTATC	AAGATCAAGG	GTACTTAGTG	CGCAATCTTT	TTGTACTGGT	2820
TTTGTTTTAT	ATACTCATCT	GTTGGCTCAG	TGCCAAACTA	ACACGGTCAG	AAAAGATAT	2880
TAAAGGCTCA	GGAATTCCCTC	AAGTCGAAGC	CGAACTGAAA	GGCCTCATGT	CCCTCAACTG	2940
GTGGGGCATT	CTTTGGAAAA	AATATGTGCT	AGGTATTCTT	GCTATTGCCA	GTGGACTCAT	3000
GCTGGGTGCA	GAGGGACCCA	GCATTCAACT	TGGAGCAGTT	GGTGGTAAAG	GAATTGCCAA	3060
GTGGCTAAA	TCCAGTCCAG	TAGAGGAACG	TTCCTTGATT	GCCAGTGGAG	CTGCAGCAGG	3120
TTTAGCCGCA	GCCTTTAATG	CTCCTATTGC	AGCACTCTC	TTTGTGTAG	AAGAAGTCTA	3180
TCACCACTTT	TCGGCCTTTT	TCTGGCTCTC	ACTCTAGCA	GCCAGCATCG	TAGCAAACCTT	3240
TGTGCTCTA	CTCATGTTCG	GTTTGACACC	AGTATTGGAT	ATGCCAGATA	ACATTCCCTCC	3300
CATGACCCCTA	GATCAGTATT	GGATATATCT	CGTCATGGGA	ATTTTCCTTG	GATTTTCAGG	3360
TTTTCTCTAT	GAGAAAAGCTG	TATTAACAGT	TGGAAGAGTT	TATGACTTGA	TTGGTCAAAA	3420
AATCCATTTG	GATAGGGCTT	ATTATCCCAT	CTTGGCTTTT	ATCCTTATCA	TACCACTCGG	3480
AATCTCTTA	CCTCAAATCA	TTGGTGGCGG	AAATCAGCTT	GTCCCTTCTT	TAACGTAAAC	3540
AAATTTAGT	TTCCAAGTTT	TATTAAGCTTA	CTTTTAATC	CGCTTTATT	GGAGTATGAT	3600
TAGCTATGGA	AGTGGACTGC	CAGGAGGAAT	TTTCCTCCCC	ATTTTAGCTC	TTGGTTCTTT	3660
GCTTGCTGCC	TTAGTTGGTG	TTATCTGTGT	CAATCTTGG	CTTGTAGTC	AAGAGCAATT	3720
CCCTATATTT	GTCATTCTAG	GAATGAGTGG	CTATTTGG	GCCATATCAA	AAGCTCCCTT	3780
AACCGCTATG	ATCCTCGTAA	CTGAGATGGT	AGGAGATATT	CGCAACCTTA	TGCCACTTGG	3840
TCTTGTCACT	CTTGTCTTT	ATATTATCAT	GGATTTGCTC	AAAGGTACGC	CAGTCTATGA	3900

257

AGCCATGCTG	AAAAAAATGC	TTCCAGAAGA	AGTATCTAGC	GAAGGAGAAG	TTACACTTAT	3960
CGAAATACCA	GTTCCTGATA	AAATTGCTGG	GAAACAAGTT	CATGAACCTCA	ACTTACCACA	4020
CAACGTCCTC	ATCACAACCTC	AAGTCCATAA	TGGCAAGAGC	CAAACAGTTA	ACGGCTCAAC	4080
CAGAATGTAT	CTGGGTGATA	TGATTCACCT	GGTTATTCCA	AAAAGTGAA	TTGGAAAAGT	4140
CAAAGATTG	TTGTTGTAGT	ATGAGTATTT	ACATAATTAA	TGTTATGTAA	ATGATCAGTT	4200
TGATTATTT	AGAAAACCGA	TTCTCAGGAA	TGAGATCGGT	TATTTTTAC	TGATGAGGAA	4260
TTTTACATAT	AAATAATTGA	ACTTTATTAA	AAATAAGACT	ATAATTAAGT	TAGAAATGAT	4320
AAAGTATAAA	GCTAGAAAGG	AGTTTACTGT	ATCAAATCTG	TACAGTAAGA	TTAAAATCAT	4380
GAAAAAGAAA	ACAATAGCAA	TTATATAGAG	AAATGAAATA	GAAATAGGAT	AAAACAATCA	4440
GGACAATCAA	ATCAATTCTC	AGCAATGTTT	TAGAAGTCCA	GATGTAACAT	TCTAGTTCA	4500
ATCTATTATA	CAATGTGTTT	TGTATCTCAT	AGCTCCTTAT	ATAGCTCTTC	AGTTATGTAG	4560
TATTAACAGA	AGTTTAGTGG	GTGAGATTTT	TATTATTTTC	CTTATTCTGT	TTTGTGTTGTA	4620
GGTCTAAGTC	TTTTTATCAC	TTTAAAAAC	TCCTATAACA	TCTTCCGAA	AAACTATAAT	4680
TTTCTTGAAA	AATATACAAG	TCTATGCTAT	ACTACTAGTA	TACTTACTTA	TGGAGAAAAT	4740
ACATGAAACG	TGAGATTTA	CTGGAACGAA	TCGACAAACT	AAAACAACTC	ATGCCCTGGT	4800
AAGTTCTGGA	ATACTACCAA					4820

(2) INFORMATION FOR SEQ ID NO: 20:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 21338 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 20:

CTACGACATC	ATGATTAACA	GTCATGGCCT	ACTACCAACT	GAGCTATGGC	GGATAAAATA	60
GTCCGTACGG	GATTCGAACC	CGTGTGTTACCG	CCGTGAAAAG	GCGGTGTCCTT	AACCCCTTGAA	120
CCAACGGACC	TTCTATCTGT	AGCAGATATA	ACCATTATAT	CAATTCTTG	CTAATTGTCA	180
ATCACTTTG	AGATTTTTC	TCTAAATAT	CTTTAATTT	TCTAATTGTTT	AATCTTGAAA	240
TAGGACAAACG	ATGGTCTTCA	TAGAAAACAA	TTTCTAAGTT	TTTCGATCA	ATTTCTCTGA	300
TATTACCTAT	ATTTACAAA	AATGACTTGT	GAGGAGAATA	AAATCGCTGA	GTATGTTGT	360
CCTTTCTG	AATATCTGTC	ATGGTACCAT	AAAACCTTTT	TGCAAAATTC	TTACCAATAA	420

258		
TGCGCAATTT	ATGAGATAACC CCTGTTGTTT CAATATACAA AATATCATGG TAAGGAATT	480
TTAAATCATT	TCCCTTGTAA TTGTAGTCGA AATAATCTAC AACATCTTCA TTTTCAAGTA	540
ACATACTCTT	CGTGTAGAAG ATATTTTGCT CAATTCTCTT CTTAAACATC TCATCATTGA	600
TATCCTTATC	AACAAAATCT AGGGCTGATA CCTGGTATTT ATAGGTTAGA GTCGCAAAC	660
CTGATCGACT	AGTGATAAAG ACGATAATAG CGTAAGGATT GTAATGACGA ATGAGCTGAG	720
CCACTTCAAA	TCCCTTTTC TCAATTCCAT GAATATCGAT ATCTAGGAAA TAAAGCTGAT	780
TTACTTCATC	ATTTCAATG TATTCTTCAA ATTCACGGAC TTTTCCCGTT GTCTTGTATG	840
ATATTGGAAT	ATTCGATTCT TTGAAATTT CATCCAATAT TCTCTCTAGT CTCACTTGAT	900
GTTCAATAAC	ATCTTCTAAA ATAAAACCTT TCATTCAAAT TCCCTCTTAA ATCTAATGAT	960
TTGTCTAAAT	GTACTGCCTT CCATCTCTGT TTCTAAAATA ATATTGTTGT ACTTATCTAG	1020
TAGTTCTTTC	ACATTATTAA ATCCGACTCC GCGATTTCTT CCCTTAGTGG AGAATCCTAA	1080
GGCAAATAGA	TCTCCTGAAG GAGTCATCGT CATTTCACAT GAATTCTGAA TCACAATAAC	1140
TGTTTCAGTT	TCCATCTTAA TAACTGCTAC TTCCATCTGC TTTTATAGC TATCAGCCGA	1200
TCCCTTGACAA	GCATTATTCA ATAAAACGCT CATGATACGA ACCAAATCCA ATAGTTCAAT	1260
TGGAAGCTTG	GTAATCGTAT CTTTTACTTC CAGTGTAAAC TCTACACCAT TATTCGAGC	1320
ATAGACAATT	GACTGAGCAA CCAAACCTCG TAAAGCTGAG TCTTCTATGT TGTTCAAATC	1380
AAAGTAAGTG	TACTTATCTG AACGCAATTT ATGATTGCT TTGACTAAAA CTTCATTTGTA	1440
AATTCTGTCA	ATTTCTGTAA AATTACCACT GTCAATTGCC ATCTGCATGC TGACAAGCAT	1500
TCCAGCATAA	TCATGTCGAA AACACCGGAT TTCATTATAC AGACCAACAA TTTCATCTGT	1560
GTAATTCTGT	AAATGTTCT GTTCAAATTT CTTCTGCTTC AAAGCAATCT CTTTCTCCAT	1620
TTGAACTTTA	TGAGAATTCA TTGCAAAGAA GGTCAAAGG AGAGAGATAA AGACAATAGA	1680
TGACAAAATA	CTTCCAAAAC TATTCACATG TTAAATCGTA CTTACCATAT CTGAAACGAA	1740
AGATACAAATA	TGTAGCAATA GTAAAGCAA AAATACCTTT TTCAAGAAAG GATAAAGGTA	1800
GTCCTTGTCA	AAATAGGCTA GTTCCAAATG GAAATAGTAA ATGATTTTTA ATGTAACAAA	1860
ATAGGTTAAC	ACCGTCACAA CGAAAAAGAA TGGGAAATGA TATTGTAAAA CAAAATTATC	1920
TCCTGTTATA	GAGGAGAAAA TTACGGACAG AAAGTTATGA GTGCTCTCAT ATAAAAGAGA	1980
TAGTAGTAAA	CTTAGGAATA GTCTCTATC CCTCTCATAC TGTTTCATCC ATCGAAAATA	2040
GGAATATAAG	CCCAAAGGAA ATAAAATCT TTCAATCCCT ATTTTATCTA AATATAGAAG	2100
ATAAAAGGAA	AATTCAAGTA CTATTTCACT TAGTAATGTA TAAGCACCAA AAACGTATAA	2160
TTCTTTCTA	TTTATTCGAC CTTTACAAAT TAAACGGTAA CTGTGACTAA TAATTAACAA	2220

259

ATGAAACAATA ACTGTCCCAA ATCCAAGTAA ATCCATTACT CTTTCTCCTT ATTCATTAC	2280
TTTTTCGTA GGAAAAGAAA ATCAAGGATG ATTCTTGAAA TCCTCATCTC CCCACCTTA	2340
ATCTTTGTA AGTCTTTTC CTTCAAAGCT ACAAACTGTT CCAATTAAAC TGTGTTTTC	2400
ATAATAAAAT CTCTAAAT GTTTTTCTT GTAAGCTAAC TTACAAAAAC CATTATACAA	2460
AATGGAATTT CGTTTAGAT AAAATTCTCT CAACTGTCAT TTTTTCTCC CAAAGTGTAC	2520
TTTTTAAGA AAAAGCCGG GAAAATTCCC AGCTTGCTA TTATATTGAT CCCAGCAGGA	2580
TTCGAACCTG CGACCGTTCG CTTAGAAGGC GAATGCTCTA TCCAGCTGAG CTATGAGACC	2640
TAATACAATT ATTCTACCAA AAATTCAATT AAAAGTCAAT TTTCTATTAA TGGTAGGGGA	2700
ATCCCTGCTG AATCGTAAA GCGCGATAGA TTTGTTCAAC AAGAACTAGT CTCATTAAC	2760
GATGGGGTAA GGTTAGGCGA CCAAAACTGA CAGAAAGATT GGCTCTATTT TTTACAGATG	2820
ATGATAATCC TAAACCTTCCC CCAATAATAA AAGTAAGAGT AGAAAATCCT TTTATAGAAG	2880
TTTCTCTAA CTGCTTACTA AATTCTCTG AGAAGAAAGT TTTCCCTTC ATGGCTAAC	2940
CAATAACGAA ATCACGGTCA GCAATTTCG ATAAAATTCT CTGACCTTCT ATTTCTAAA	3000
TCTTTGATT TTCTGATTCA CTGGCCTTAT CTGGTGTGTT TTCATCTGAT AACTCAATCA	3060
TTTCAAACCTT AGCAAATCTA GAAATTCTG TTGAATACTC TGCGATACCA TCTTTAAAT	3120
ACTTTCTTT CAGTTCCCA ACTGTTACAA CTTTAATTTC CATGACTCTA TTCTAACATA	3180
TTCTCTATTT TTTCACATCT TATTCAACAA ATAAAAAAATA GATTCAATT AAGAAAAATCA	3240
CAATTCAAA AGAGTTATCC ACAGTTGTG TAAAACCTTT GTGTTAAGT TATAATTAG	3300
CTAGTCAGTT TATACCTTCA GTAATTCAA CATATGGAGG CAAATATGAA ACATCTAAA	3360
ACATTCTACA AAAATGGTT TCAATTATTA GTCGTTATCG TCATTAGCTT TTTTAGTGGA	3420
GCCTTGGGTA GTTTTCAAT AACTCAACTA ACTCAAAAAA GTAGTGTAAA CAACTCTAAC	3480
AACAATAGTA CTATTACACA AACTGCCTAT AAGAACGAAA ATTCAACAAAC ACAGGCTGTT	3540
AACAAAGTAA AAGATGCTGT TGTTCTGTT ATTACTTATT CGGCAACAG ACAAAATAGC	3600
GTATTGCGA ATGATGATAC TGACACAGAT TCTCAGCGAA TCTCTAGTGA AGGATCTGGA	3660
GTTATTCTATA AAAAGAATGA TAAAGAAGCT TACATCGTCA CCAACAATCA CGTTATTAAT	3720
GGCGCCAgCA AAGTAGATAT TCGATTGTCA GATGGGACTA AAGTACCTGG AGAAATTGTC	3780
GGAGCTGACA CTTCTCTGA TATTGCTGTC GTCAAAATCT CTTCAGAAAA AGTGACAACA	3840
GTAGCTGAGT TTGGTGATTC TAGTAAGTTA ACTGTAGGAG AAACCTGCTAT TGCCATCGGT	3900
AGCCCGTTAG GTTCTGAATA TGCAAATACT GTCACTCAAG GTATCGTATC CAGTCTCAAT	3960

260

AGAAATGTAT CCTTAAAATC GGAAGATGGA CAAGCTATTCTACAAAAGC CATCCAAACT	4020
GATACTGCTA TTAACCCAGG TAACTCTGGC GGCCCCTGTA TCAATATTCA AGGGCAGGTT	4080
ATCGGAATTA CCTCAAGTAA AATTGCTACA AATGGAGGAA CATCTGTAGA AGGTCTTGGT	4140
TTCGCAATTCTGCAATGA TGCTATCAAT ATTATTGAAC AGTTAGAAAA AAACGGAAAA	4200
GTGACGCGTC CAGCTTGGG AATCCAGATG GTTAATTATCTAATGTGAG TACAAGCGAC	4260
ATCAGAAGAC TCAATATTCC AAGTAATGTT ACATCTGGT TAATTGTTCG TTCCGGTACAA	4320
AGTAATATGC CTGCCAATGG TCACCTTGAA AAATACGATG TAATTACAAA AGTAGATGAC	4380
AAAGAGATTG CTTCATCAAC AGACTTACAA AGTGCTCTT ACAACCATTC TATCGGAGAC	4440
ACCATTAAGA TAACCTACTA TCGTAACGGG AAAGAAGAAA CTACCTCTAT CAAACTTAAC	4500
AAGAGTTCAAGTGTAG ATCTTAATTG ACATCTATGT AAAGAAAGCT TTACATAAGA	4560
GAAAAGATGT GTTAGTGTAG AATCATGGAA AAATTTGAAA TGATTTCTAT CACAGATATA	4620
CAAAAGATC CCTATCAACC CCGAAAAGAA TTGATAGAG AAAAAGCTAGA TGAACTAGCA	4680
CAGTCTATCA AAGAAAATGG GGTCAATTCAA CCGATTATTG TPCGTCAATC TCCTGTTATT	4740
GGTTATGAAA TCCTTGCAGG AGAGAGACGC TATCGGGCTT CACTTTAGC TGGTCTACGG	4800
TCTATCCAG CTGTTGTTAA ACAGATTCA GACCAAGAGA TGATGGTCCA GTCCATTATT	4860
GAAAATTTAC AGAGAGAAAA TTAAACCCCA ATAGAAGAAG CACGCGCCTA TGAATCTCTC	4920
GTAGAGAAAG GATTCAACCA TGCTGAAATT GCAGATAAGA TGGGCAAGTC TCGTCCATAT	4980
ATCAGCAACT CCATTCTGTTT ACTTTCCTTG CCAGAACAGA TTCTTCAGA AGTAGAAAAT	5040
GGCAAACCTAT CACAAGCCCA TGCGCGTTCC CTAGTTGGGT TAAATAAGGA ACAACAAAGAC	5100
TATTTCTTTC AACGGATTAT AGAAGAAGAT ATTTCTGTAA GGAAATTAGA AGCTCTTCTG	5160
ACAGAGAAAA AACAAAGAA ACAGCAAAAA ACTAATCATT TCATACAAAA TGAAGAAAAAA	5220
CAGTTAAGAA AACTACTCGG ATTAGATGTA GAAATTAAAC TATCTAAAAA AGACAGTGGAA	5280
AAAATCATTA TTTCCTTTTC AAATCAAGAA GAATATAGTA GAATTATCAA CAGCCTGAAA	5340
TAAGGCTGTT CTTTATTTT TTTATCTCAC AAGGTTATCC ACTATGTTTT TCGATAAAAA	5400
GCTTAATAAA TCAATAATTCTTCTTTAT CCCAACCTG TGGATAAAAGT TTGGTAACAT	5460
TGTGGATTAT TTTCACAGC TTGTGGAAAA TTCTTGCTAT CTATGGTAAATATCTCTAG	5520
TATTAACCTT TTAAATAGTA AAGGAGGAGA AAGGATTGAA AGAAAACAA TTTTGGAAATC	5580
GTATATTAGA ATTTGCACAA GAAAGACTGA CTCGATCCAT GTATGATTTC TATGCTATT	5640
AAGCTGAACCTACATCAAGGTA GAGGAAATG TTGCCACTAT ATTTCTACCT CGCTCTGAAA	5700
TGGAAATGGT CTGGGAAAAA CAACTAAAAG ATATTATTGT AGTAGCTGGT TTTGAAATTT	5760

261

ATGACCGCTGA	AATAACTCCC	CACTATATTT	TCACCAAACC	TCAAGATACG	ACTAGCTCAC	5820
AAGTTGAAGA	AGCTACAAAT	TTAACTCTTT	ATAACTATAG	TCCAAAGTTA	GTATCTATT	5880
CTTATTCAAGA	TACGGGATTAA	AAAGAAAAGT	ATACCTTGA	TAACTTTATT	CAAGGGGATG	5940
GAAATGTTG	GGCTGTATCA	GCCGCTTTAG	CTGTCTCTGA	AGATTTGGCT	CTGACCTATA	6000
ACCCCTTTT	TATCTATGGA	GGACCAGGCC	TTGGTAAGAC	TCACTTATTA	AACGCTATTG	6060
GAAATGAAAT	TCTAAAAAAAT	ATTCTAATG	CCCGTGTAA	ATATATCCCT	GCCGAAAGCT	6120
TTATTAATGA	CTTTCTTGAT	CACCTAAGAC	TTGGGGAAAT	GGAAAAGTTT	AAAAAGACCT	6180
ATCGTAGTCT	TGATCTTTG	TTAATCGATG	ATATCCAGTC	ACTCAGCGGA	AAAAAAGTCG	6240
CAACTCAGGA	AGAATTTTC	AATACCTTCA	ACGCCCTTCA	TGACAAGCAA	AAACAGATTG	6300
TCCTAACGAG	TGATCGTAGT	CCAAAACATC	TAGAAGGGCT	CGAGGAGAGG	CTTGTACCGC	6360
GTTTTAGTTG	GGGATTGACA	CAAACATATCA	CCCCCCCTGA	CTTGAAACA	CGTATTGCCA	6420
TTTTACAAG	TAAGACGGAA	CATTTAGGCT	ACAATTTC	AAGTGATACT	CTAGAATACC	6480
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CCAAAATAAA	ATCTTGATT	GATCAAGACG	ATAATTACG	TTTAGAAATT	GAATCAATCA	6900
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TGAAGATGCT	GGTTTGTAA	TTACTTCTTT	AGGTTCGATC	CTTCTTGAAG	CTTCTTCTT	7320
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262	
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264

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266

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268

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ATGAATATTA	CCTTCATTAA	TGGAAATTGT	CGGATTGGGA	GCAAGAAGGC	ATGCTCTGCT	19920

TACATGAATT GATTAGTAGA GAAGAAGGAC TGGTAGACGA TATTCCACGT TTAAGGAAAT	19980
ATTTCAAGAC CAAGTTCGA AATCGAATTT TAGACTATAT CCGTAAACAG GAAAGTCAGA	20040
AGCGTAGATA CGATAAAAGAA CCCTATGAAG AAGTGGGTGA GATCAGTCAT CGTATAAGTG	20100
AGGGGGTCT CTGGCTAGAT GATTATTATC TCTTTCATGA AACACTAAGA GATTATAGAA	20160
ACAAACAAAG TAAAGACAAA CAAGAAGAAC TAGAACCGCT CTTAAGCAAT GAACGATTTC	20220
GAGGGCGTCA AAGAGTATTA AGAGACTTAC GCATTGTGTT TAAGGAGTTT ACTATCCGTA	20280
CCCACCTAGTA AGTCATGCAA AAAAAATGAA AAAAATTAGA AAAAGTAGTT GACAAAGTTT	20340
GAAAAGGCTG TATAATAGTA AGAGTTGAAA ATAACAACTC AGGTCCGTT GTCAAGGGT	20400
TAAGACACCG CCTTTTCACG GCGGTAACAC GGTTCGAAT CCCGTACGGA CTATGGTATG	20460
TTGCGTCAGG ACCACTTGAT GAAAAAAAGT TAAAAAAAC TAAAAAAATCT TCAAAAAAAGT	20520
GTTGACAAGC GAAAGCAGTT GTGATATACT AATATAGTTG TCGCTTGAGA GAAGCAAGTG	20580
ACAAAGACCT TTGAAAACGT AACAAAGACGA ACCAATGTGC AGGGCGCTAC AACGTAAGTT	20640
GTAGTACTGA ACAATGAAA AAACAATAAA TCTGTCAGTG ACAGAAATGA GAAAGCAACTC	20700
AAACTTTTA ATGAGAGTTT GATCCTGGCT CAGGACGAAC GCTGGCGCG TGCTTAATAC	20760
ATGCAAGTAG AACGCTGAAG GAGGAGCTTG CTTCTCTGGA TGAGTTGCGA ACGGGTGAGT	20820
AACCGTAGG TAACCTGCCT GGTAGCGGG GATAACTATT GGAAACGATA GCTAATACCG	20880
CATAAGAGTA GATGTTGCAT GACATTTGCT TAAAAGGTGC ACTTGCATCA CTACCAAGATG	20940
GACCTGCGTT GTATTAGCTA GTTGGTGGGG TAACGGCTCA CCAAGGCGAC GATAACATAGC	21000
CGACCTGAGA GGGTGATCGG CCACACTGGG ACTGAGACAC GGCCCAGACT CCTACGGAG	21060
GCAGCAGTAG GGAATCTTCG GCAATGGACG GAAGTCTGAC CGAGCAACGC CGCGTGAGTG	21120
AAGAAGGTTT TCGGATCGTA AAGCTCTGTT GTAAGAGAAC AACGAGTGTG AGAGTGGAAA	21180
GTTCACACTG TGACGGTATC TTACCAAGAAA GGGACGGCTA ACTACGTGCC AGCAGCCGCG	21240
GTAATACGTA GGTCCCGAGC GTTGTCCGGA TTTATTGGC GTAAAGCGAG CGCAGGCGGT	21300
TAGATAAGTC TGAAGTTAAA GGCTGTGGCT TAACCATA	21338

(2) INFORMATION FOR SEQ ID NO: 21:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6273 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

270

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 21:

TGTTTTAAA GACCGTGTC TGGATAGACT TTCGGACGCA ACGCTCTATT AGATAATGAA	60
CTGCCTATAC ACAAGATTTC TAACTTAGT CGACATGAGC TGAAACCTCT TATTGTTAA	120
GTAGTCACA AAATATTATA CACCTATTT ATGAATAGTC AACTGTCTT ACAGTAAAAT	180
TTTAGAAAAT CATGAAAATT TTCTCTTCT TTCCATTAA AGTACATTC AGTCATTCTC	240
ACATCAAAA AGCCAGACG AAATTGTCTG AGCATTCTT TATCTAGTCG TTTAAGGAAG	300
TTGAGTCAG TATGTTAAA GTCTCTGTC CATCATTCTC TCAACAAACC TTGTTCTGG	360
AGAAACTCCT TGGCTACTTG CTTGCTGAC TTGCTTCAA CACCGACTTG GTAGTTGAGC	420
TGGCTCATCT GGCTTCTGT AATCTTACCA GCCAATGTAT TAAGAACTCT TTCCAACCTCT	480
GGGTGTTCT TGAGAAAGAGC TTCTTCATG AGTGGAGCCC CTTGATAAGG TGGGAAGAGT	540
TGCTTGTCAT CTTCCAAGAC CTGTAATCA TAAACGCTCAA ATTCCGCATC AGTCGAATAG	600
GCATCCGTGA TTTGAATATC CCCTGACTGA ATAGCCTGAT AGCGAAGGGC TGGCTCAATG	660
GTCGCTACAT TGAGATTGAG ACCATACATT GATTGCAAGC CCTTATTTC ATCTTCACGG	720
TCGTTAAACT CGAGTGTAAA ACCTGCCTTC AACTGCCCTT CCACTTTTT CAAGTCTGAA	780
ATGGTCTTCA AGCCATATT TTGAGCAATC TTTTCGGAA CAGCTACAGC ATAGGTGTTT	840
TGATAAGACA TGGGTTTGAG ATAGGCTAGA TGATCCTGCT TAGCAATGCC ATCACGCGCC	900
ACCTGATAAA CCTGTTCTGG TTCATGACTC ACCTGGGTG ATGGTTGAAG CAAACTTTCA	960
GTCACCGTAC CAGTAATTC AGGATAGATG TCAATATCGC CTTTTTCAG AGCTTCATAA	1020
AGGAAGCTTG TCTTCCAAA ATTCCGTTA ACAGTCGGAG TCATGCTGGT ATTTTCTTCA	1080
ATCAGCAACT TATACATATT GCCAAAATT TCTGGTTCTG GACCTATTTC CCCAGCAATA	1140
ACCAAGTTT CCTTCTCTT TTGAACCAAAGAGCTGGAC TATAAGACAG ACCCAGTAAT	1200
AAAGCCACCA AGGCAAAACC TGAGAAAATC GTCCGTAATT TTGCTTTTC CATCACTTTT	1260
AGTAGGAAGT TAAAGGCAAT GGCTAGCACT GCAGAAGAAA GTGCCCAAT CAAAATCAA	1320
CTGGCATTAT TACGGTCAAT TCCAAAAGA ATAAAGGAAC CTAGTCCCC TGACCAATC	1380
AAGGCCGCCA AGGTTGCCGT ACCGATAATC AAAACAGCTG CCGTCCGAAT CCCAGACATG	1440
ATAACAGGCA TGGCGAGTGG AATTCAAT TTCTTGAGAC GTTCCCATCT GGTCATCCCA	1500
AAGGCAATCC CAGCCTCTTG CAGGTTCGGA TCAATTCCCT TCAGCCCAAGT GATAGTATT	1560
TGCAAAATAG GGAAAATCGC ATAAATCACT AGAGCTGTCA AAGCCGGCAA GGTCCCAATT	1620
CCCATCAAAG GGATAAAAGAG CCCCAACAAG GCCAGAGACG GGATGGTCTG GAAAATACCT	1680
GCAATCTGCA AGACCCAGTC GGCCAGCTTC TCATGATAGC GAAGAAAAAC AGCCAAGGGA	1740

ATCGCAAGCA AAATAGCTAG TAACAAGGTC AAAAGCGACA ACTGCAAATG TTGAGATAGA	1800
GCTGTCACC AATCACTAAA ACGATCCTGA AAAGTTGCAA TTAAATTAGT CATGAACACT	1860
ACCTCCAAAC AAGTCTGCTA CAAAGTCTGT TGCAGGCCTGCT TTTAAAATTG TCTCGGGATT	1920
CGCTACCTGG CGAATTTCTC CATCCTGCAA GACAGCAATA CGGTCCGCCA ACTTCAGGC	1980
TTCATCCGTA TCATGGGTTA CAAAATCGT TGTCATCCCA AACTCTTTAT GCAATTCTTT	2040
TGTCAGAACC TGCAACTGTT TTCTCGAAAT AGCATCCAAG GCCGAAAAGG GTTCATCCAT	2100
GAGGAAAATC TTGGGCTGAC CAATCATAGC TCGGACAATA CCGACCCGTT GCTGTTCTCC	2160
ACCAGATAAT TCACTAGGTA AGCGATGCCC ATACTCGGCT ACTGGTAAAC CAACCTTAGC	2220
CAAAAGCTCT TCTGTTTCTC TCGTAATTTC TTCCCTGCTC CACCCCTTC ATTTCAGGAAT	2280
GAGAGCAATA TTTCCGCAA CTGTTAGATT TGGAAAAAGA GCAATAGCCT GTAAAACATA	2340
ACCAGTAGAA AGACGAAGTT CACGCTCATC ATAGTCTTTG ATGCGCTTCC CATCCATATA	2400
AATATTTCCA TCAGTTGGTT CAAAAGACG GTTAATCATC TTGAGCATGG TCGTCTTACC	2460
TGACCCAGAA GGCCCTACTA AAACCATAAA TTCCCCATCC TCAATCTGTA AGTTGACATC	2520
TCTCAAGACA TCCTTTCTG TGTAGCGCAG TGCTACATTT TTGTATTCAA TCATTCTTTG	2580
TCCTCAATTT AAAACTTCCC TCGATTGGTC AAGTCTTCTA CCTTAGGCAT AACTCCCTTA	2640
TTATCCCAAT GCTCCACAAT TTTCGGTTC TCTAAACGGA AGATATCGTA CTGGCATAA	2700
GCAACGCCAT CAATCTGAGT CTGACCCATAC CTAACCACAT AGTTCCCTTG TCCTAAAGAGT	2760
TGGAAAACAA AGTCAAAAGT GACACTATAT TCAGCCACAT AGTTTTTATA AGCAGCACTT	2820
CCTTGTCCAA TATCATGATT ATGCTGAATC AAATCGTCTG CCACATAATC ACTCCACTGC	2880
TCTAGCTCCC CATTGGAA AATTCTGTC AAGAAACGGC GAACCAGCTT TTATTTCT	2940
GCTTTCTTAT CCAAATCCTT GATTCAAAA TCTCCAAAAA TTGATCTAG TTGGTCATT	3000
TCAGGTGTT GATAGTAGTC AATGACATCC CAATGCTCAA CAATACAACC ATTCTCATCC	3060
TCACGGAAAG TATCCGTCGT CACCCATTGA GCTTCTCCAC CATTAGATA TTGATGAACA	3120
TGAACAAAGA CCAGATTGCC ATCCCTCAATG GTGCGGACAA TCTTAATCTG ACGCTCTGGA	3180
TGACGCTCAA AGAAATCTGC AAAGAAGGCT GCAAATCCTT CTTTCCCGTC AGGAACACCT	3240
GTCGAATGTT GGATATAGGT ATCCCCTACA GACTGGGCTT GAGCCTCAGC AACTCGTCCG	3300
TCTTGAATGG CATGGATGTA TAGGTTGTGA GCATTTTCA CTTGTTGTGA CATATTCTAA	3360
ACCTCATTTC CCTTCTCTTT CAGATTGCC AAAATTCTTT CTTGAAAACC TTCAAATTGG	3420
TGAATTCTT CCTCTGAAA TCCTTTGTAA AAGATAGTAT CCAATTCTG ACTGACACGA	3480

272	
TGCCCCACTT	3540
CTTTCTGGGA	
CTTGCCTAAC	
TCCGTTAAAA	
CTAAATACTT	
CTTACGCTTG	
TCTTTTCCAC	3600
ACGGACTAAC	
AATTACAAGC	
TTTTGTTCCCT	
CTAGCTTTTT	
TATCATAGTC	
GTCAGCGTAT	3660
TATTGCGAAG	
TCCAGTCGCA	
AGCGCGATAT	
CTGTCGCAGT	
TGCGCAGCCA	
GTTCACTAT	3720
TCCATAAAAC	
CGCTAAAATC	
TTGCCCTGTT	
CACCCCTATA	
AAGAGCCTCA	
GGATCTTGAC	3780
TCAGTAACCTT	
TTGAAAATC	
CGCCCATTCA	
ACAAACGAAT	
ATGATGGGCT	
AGCAAATGAC	3840
CATCTTCAT	
AACACCTCCA	
ATTTATTCG	
ATATCGAAAT	
GAATAAAACA	
ATTGTAACAC	3900
TCATCGTTCT	
AACTGTCAAC	
TATTCGATT	
TAGAAATAAT	
TTTTGATAAT	
TATCCACACC	3960
ACCATACTCC	
GGCTCAACTA	
ACTTTAACG	
AGAGTTCTA	
AACTCCTTCG	
TCCTCCAGTC	4020
TACAAAAGCC	
TTCCATTCTG	
ACTATCCTAT	
ATTTTATGAG	
GGGACACATT	
TTTCTTATCA	4080
GACCATTAT	
TTTAAAGATA	
GAAGTAAATC	
ATAATTGCTT	
CCATCTGTT	
TTTTATAGTA	4140
TATTGAAGTT	
AGACTAGAGC	
ACTGTATCTT	
CTAAACATT	
GATAGAAAGC	
GATTTGAATT	4200
TCCCAATCAA	
TTTGTTCGTA	
TTTATAGCAT	
TTCGAAACTG	
GAATAGGACA	
CCATGACTGC	4260
TAAAAGATT	
CTATAAATTC	
ATTTAATTT	
CTCAATCAAT	
TTGTTCATAT	
CTTATTCAT	4320
TCCGCTATAA	
TTTCACCTTA	
CCCTATCTT	
TTCGTAGCAC	
CCTTCAAACA	
GCCTATCCCC	4380
TACCGTTGA	
CGATTCCCTA	
CTTCGCTCCA	
CTTCCATTAC	
AGAAGTTCT	
TCACTACTAT	4440
GGGCTCGGCT	
GACTTCTCAT	
GATTCTCTGT	
TACTACTATT	
TGAACGCTCA	
CGAGATAGAT	4500
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ATGCTTTGAT	
CCACAATGGA	
ATCAAAGCAT	
TTTAAAGAGT	
TCCTCATACA	4560
TAAGCGCAGA	
AGTCGCAGTT	
CCTCTGTACT	
TGGCTTCTTC	
TCTTTTGACA	
AAGCGAGCCA	4620
AGTTGAGCAA	
CTCAGGTGCT	
GGATGTTGG	
GATTTAGGAG	
CAATTCAACGA	
TTGACCAGGC	4680
CTGAGAGAGC	
AACTGCTTC	
ATTGCTCAT	
TTGAGTAGGAG	
CAGTTTTTA	
GTAGTCTCTA	4740
GGAGAGCAGC	
AACTAAATCT	
TCACTCAAAT	
CATGTCGAGC	
ATGATTGTAA	
AGATCTTTA	4800
TAAGGCTTTC	
TAGGTTGGT	
TCTACCATCC	
CTACCACCTC	
CCTTATGGTT	
TAATAATGTT	4860
TAATCAAATC	
AACCGTTGAA	
CGATCCAATT	
TCTTCACCAA	
GGCTTGTAAG	
AAAGCTTGC	4920
CTTCTAGGAA	
GTCATCCATT	
GCATAGAGGG	
TTTGGTGAGA	
ATGGATATAA	
CGAGCGCAGA	4980
CACCGATAGT	
TGTTGATGGG	
ACACCACCAT	
TTTCAGATG	
AGCTGCACCT	
GCATCTGTT	5040
CGCTTTTAC	
ACAGTAGTAT	
TGGTACTTGA	
TACCACTTC	
TTCAGCCGTT	
GTCAAAAGGA	5100
AATCCCTCAT	
CCCTGGGAGA	
AGCAAGTGAC	
CTGGATCATA	
GAAACGAATC	
AAGGTTCCAT	5160
CTCCAATCTT	
GCCTTGACCA	
CCGTAGACAT	
CACCTGCTGG	
TGAGCAATCA	
ACTGCGAGGA	5220
AGACTTCTGG	
GTCAAACCTG	
GTTGTAGAGG	
TATGAGCGCC	
ACGCAGACCA	
ACTTCCTCTT	5280
GGACGTTAGA	
ACCCAGATAG	
AGTTCATTCG	
CGAGTTTTTG	
ACCCGATAAA	

273

GCTTCAGCTA GCTCGCTTAC CATGAGGACA CCGTAGCGGT TATCCAAGC TTTTGAGATG	5340
ATATTTTTT CATTGGCTGT CAAAATTGCA GAACTATCTG GTACAATGGT ATCACCAGGA	5400
CGGATGCCAA AACTTTCTGC CTCAGCCTTG TCCGCAAAAC CACCATCAA AACGATATCG	5460
GCAATGGCTG GCATGGTTGG TCCCCCCTTT CCACGAGTCA AATGCGGAGG AACAGAACCT	5520
GAAATCACAG GAATTCATG ACCATCACGA GTCAAGAGTT TGAAACGTTG GCTGCTAAC	5580
ACCATGGGGT TCCAGCCACC GATTCTACG ACACGGAAGG TACCATCTGG CTTGATTTCG	5640
CTGACCATAA AACCAACTTC GTCCATATGA GAAGCGACCA AGACGCGCGG TGCATCCACA	5700
GCTTCTGAAT GTTGATACC AAAAATACCA CCCAAGCCAT CTGTCACCAC TTCATCCACA	5760
TGGGGTGTCA ACTTTTCACG AAGATAAGCA CGGACAGGCG CTTCATGACC TGAGACTGCA	5820
GCAAGTTCTG TTACTTCTTT AATTTTGAA AATAATGTTG TCATTTCACT TCCTTCTTC	5880
TTTCATCCAT TTTACCACTT TTTATAGGAG AAGGATAGTG GGAAGGTGGA TTTCTAAGTT	5940
AGTATCTTAG TCCTGCTCTA TCTTAGAAAA GGATAGTATT CTCTTGATG TAGTGCAAAA	6000
TCTAGAAAC ATTCCAAAAT TAATCGAAT ATTTATTTCC AAACAAAAAA ACAATACACC	6060
ATCAAAGTTG TTTGGATTTT TCATGAAATT TACAGAAAAT AGTTGACTTC CCTTTCTTC	6120
TTCTTTAAAT ATATAGTTGG TTGAGTTTG AATAGTACGC TGTAGCTGCT AAAACATTTC	6180
TAGAAATTAA TTTGACTTTC CTAATAGAGT TGTTCATATC TTATTTCAAT TTACTATAGT	6240
ACAAAAACTAG AAAAGGAAAA AATCATGACC AGG	6273

(2) INFORMATION FOR SEQ ID NO: 22:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 28171 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 22:

ACAACCTTTT TCAAAAATC ACCTTGGTAC GGAGATGTT TGCTTTCTGC TATTATTTTC	60
GGTTATATTC ATATCAATT TGCTTTAACT CCTCTTGCTT TTTTCATTAA TGCTAGTGGAA	120
GGTCTTATTT TAGCTCTATT GTATCGCATG ACTAAAAATC TCTACTATCC AATACTAGTT	180
CATATTCTCA TTAATATCAC TGCTTCTGG GATGTGTGGT TGCTCCTATT TTCAGGAAGT	240
TAGCTTACTA AAATAATGTC GGAACCTTCC GGCATTTCT TTTTCACAA ATAGTCAACG	300
TTTTCTTTT CGATATTGTA GTGGTGTGTA TCCAGTTATT TTTTGAAATT GATTTGAAA	360

274						
ATAAAGGTTGA	CTTGAGAAAG	GCAGATAGTG	AAGATAGTTA	AGAAGAATAG	GATGTTCTT	420
TTTCCTTTTT	GGAAAACCTTC	TAAAATATGG	TATAATGAAA	AGATAAAGAA	GTTGGGGTA	480
GAAGATGAAC	ATTCACAAAT	TACGCTATGT	TGTGGCTATT	GCCAATAGTG	GTACTTTCG	540
TGAAGCTGCT	AAAAAGATGT	ATGTTAGTCA	GCCGAGTCTG	TCTATTTCTG	TTCGTGATT	600
GGAAAAAGAG	TTGGGCTTTA	AGATTTCCG	TCGGACCAGC	TCAGGGACTT	TCTTGACCCG	660
TCGTGGGATG	GAATTTATG	AAAAATCGCA	AGAATTGGTT	AAAGGATTG	ATATTTTCA	720
AAATCAGTAT	GCCAATCCTG	AAGAAGAAAA	AGATGAATT	TCTGTTGCTA	GCCAGCACTA	780
TGACTTCTTG	CCACCAACTA	TTACGGCCTT	TTCAGAGCGC	TATCCTGACT	ATAAGAACCT	840
CCGTATTTTT	GAATCAACTA	CTGTTCAAAT	ATTAGATGAA	GTGGCGCAAG	GGCATAGTGA	900
GATTGGGATT	ATCTACCTCA	ACAATCAAA	AAAAAAGGGG	ATTATGCAAC	GGGTTGAAA	960
ATTAGGTCTG	GAGGTACATCG	AATTGATTCC	TTTCCATACC	CATATTATC	TCCGTGAGGG	1020
TCATCCTTTA	GCCCAGAAAG	AGGAATTAGT	CATGGAGGAT	TTAGCGGATT	TACCAACGGT	1080
TCGTTCACT	CAAGAGAAAG	ACGAGTACCT	TTATTATTCA	GAGAACTTTG	TCGATACCAAG	1140
CGCTAGCTCA	CAGATGTTA	ATGTGACAGA	CCGTGCCACC	TTGAATGGTA	TTTTGGAGCG	1200
GACGGACGCC	TATGGCAGACAG	GTTCTGGATT	TTTAGATAGT	GACAGTGT	ATGGCATTAC	1260
AGTTATTCGT	CTCAAGGATA	ACCTAGATAA	CCGCATGGTC	TATGTTAAC	GTGAAGAAGT	1320
GGAGCTTAGT	CAAGCTGGGA	CTCTCTCGT	AGAAGTCATG	CAAGAATATT	TTGATCAAAA	1380
GAGGAAATCA	TGAAAAAAAG	AGCAATAGTG	GCAGTCATTG	TACTGCTTT	GATTGGGCTG	1440
GATCAGTTGG	TCAAATCCTA	TATCGTCCAG	CAGATTCCAC	TGGGTGAAGT	GGCCTCCTGG	1500
ATCCCCAATT	TCGTTAGCTT	GACCTACCTG	CAAATCGAG	GTGCAGCCTT	TTCTATCTTA	1560
CAAGATCAGC	AGCTGTTATT	CGCTGTCATT	ACTCTGGTTG	TCGTGATAGG	TGCCATTGG	1620
TATTACATA	AACACATGGA	GGACTCATTC	TGGATGGCT	TGGGTTGAC	TCTAATAATC	1680
CGGGGTGGTC	TTGGAAACTT	TATTGACAGG	GTCAGTCAGG	GCTTTGTTGT	GGATATGTT	1740
CACCTTGACT	TTATCAACTT	TGCAATTTC	AATGTGGCAG	ATAGCTATCT	GACGGTTGGA	1800
GTGATTATTT	TATTGATTGC	AATGCTAAA	GAGGAATAA	ATGGAAATTA	AAATTGAAAC	1860
TGGTGGTCTG	CGTTGGATA	AGGCTTGTC	AGATTGTC	GAATTATCAC	GTAGTCTCGC	1920
GAATGAACAA	ATTAATCAG	GCCAGGTCTT	GGTCAATGGT	CAAGTCAGA	AAGCTAAATA	1980
CACAGTCCAA	GAGGGTGATG	TCGTCACCTA	CCATGTGCCA	GAACCAGAGG	TATTAGAGTA	2040
TGTGGCTGAG	GATCTTCCGC	TAGAAATAGT	CTACCAAGAT	GAGGATGTGG	CTGTCGTTAA	2100
CAAACCTCAG	GGAATGGTTG	TGCACCCGAG	TGCTGGTCAT	ACCAAGTGGAA	CCCTAGTAAA	2160

TGCCCTCATG TATCATATTA AGGACTTGTC GGGTATCAAT	GGGGTTCTGC GTCCAGGGAT	2220
TGTTCACCGT ATTGATAAGG ATACGTCAGG TCTTCTCATG	ATTGCTAAA ACGATGATGC	2280
GCATCTAGCA CTTGCCAAG AACTCAAGGA TAAAAAGTCT	CTCCGCAAAT ATTGGCGAT	2340
TGTTCATGGA AATCTACCTA ATGATCGTGG TGTAAATTGAA	GCGCCGATTC GCCGGAGTGA	2400
AAAAGACCGT AAGAACACAGG CTGTAACCTGC TAAAGGGAAG	CCTGCAGTGA CGCGTTTCA	2460
CGTCTTGGAA CGCTTGGCG ATTATAGCTT AGTAGAGTTG	CAACTGGAGA CAGGGCGCAC	2520
TCATCAAATC CGTGTCCACA TGGCTTATAT CGGCCATCCA	GTCGCTGGTG ATGAGGTCTA	2580
TGGTCCTCGC AAGACTTGA AAGGACATGG ACAATTCTT	CATGCCAAGA CTTTAGGTTT	2640
TACTCATCCG AGAACAGGTA AGACCTTGGA ATTAAAGCA	GATATCCAG AGATTTTAA	2700
GGAAACCTTG GAGAGATTGA GAAAGTAAGA ATGAAAAGA	ATTAACCTAG TTTAGCACTT	2760
GTAGGCGCTT TTTAGGTTT GTCATGGTAT GGGAAATGTT	AGGCTCAAGA AAGTCAGGA	2820
AATAAAATCC ACTTTATCAA TGTCAAGAA GGTGGCAGTG	ATGCGATTAT TCTTGAAAGC	2880
AATGGACATT TTGCCATGGT GGATACAGGA GAAGATTATG	ATTTCAGAAGA TGGAAAGTGT	2940
TCTCGCTATC CATGGAGAGA AGGAATTGAA ACGTCTTATA	AGCATGTTCT AACAGACCGT	3000
GTCTTCGTC GTTGAAGGA ATTGGGTGTC CAAAAACTTG	ATTTTATTTT GGTGACCCAT	3060
ACCCACAGTG ATCATATTGG AAATGTTGAT GAATTACTGT	CTACCTATCC AGTTGACCGA	3120
GTCTATCTTA AGAAATATAG TGATAGCTCT ATTACTAATT	CTGAAACGTCT ATGGGATAAT	3180
CTGTATGGCT ATGATAAGGT TTTACAGACT GCTGCAGAAA	AAAGGTGTTTC AGTTATTCAA	3240
AATATCACAC AAGGGGATGC TCATTTTCAG TTTGGGACA	TGGATATTCA GCTCTATAAT	3300
TATGAAAATG AAACGTGATTC ATCGGGTGAA TTAAAGAAAA	TTTGGGATGA CAATTCCAAT	3360
TCCTTGATTA GCGTGGTGAA AGTCAATGGC AAGAAAATT	ACCTTGGGG CGATTTAGAT	3420
AATGTTCATG GAGCAGAAGA CAAGTATGGT CCTCTCATTG	GAAAAGTTGA TTTGATGAAG	3480
TTTAATCATC ACCATGATAC CAACAAATCA AATACCAAGG	ATTTCATTAA AAATTTGAGT	3540
CCGAGTTGTA TTGTTCAAAC TTGGATAGT CTACCTTGGA	AAAATGGTGT TGATAGTGAG	3600
TATGTTAATT GGCTCAAAGA ACCAGGAATT GAGAGAATCA	ACCGCAGCCAG CAAAGACTAT	3660
GATGCAACAG TTTTGATAT TCGAAAAGAC GGTTTGTCA	ATATTCACAC ATCCTACAAG	3720
CCGATTCAA GTTTCAAGC TGTTGGCAT AAGAGTGCAT	ATGGGAAC TGTTGATCAA	3780
GCGCCTGATT CTACAGGAGA GTATGCTGTC GGTTGGAATG	AAATCGAAGG TGAATGGTAT	3840
TACTTTAACCC AACCGGGTAT CTGTTACAG AATCAATGGA	AAAAATGGAA CAATCATTGG	3900

276	
TTCTATTTGA CAGACTCTGG TGCTTCTGCT AAAATTGGA AGAAAATCGC TGGAATCTGG	3960
TATTATTTA ACAAAAGAAA CCAGATGGAA ATTGGTTGGA TTCAAGATAA AGAGCAGTGG	4020
TATTATTTGG ATGTTGATGG TTCTATGAAG ACAGGATGGC TTCAATATAT GGGGCAATGG	4080
TATTACTTTG CTCCATCAGG GGAAATGAAA ATGGGCTGGG TAAAAGATAA AGAAACCTGG	4140
TACTATATGG ATTCTACTGG TCTCATGAAG ACAGGTGAGA TAGAAGTTGC TGGTCAACAT	4200
TATTATCTGG AAGATTCAGG AGCTATGAAG CAAGGCTGGC ATAAAAAGGC AAATGATTGG	4260
TATTTCTACA AGACAGACGG TTCACGAGCT GTGGGTTGGA TCAAGGACAA GGATAAAATGG	4320
TACTTCTTGA AAGAAAATGG TCAATTACTT GTGAACGGTA AGACACCAGA AGTTTATACT	4380
GTGGATTCAA GTGGTGCCTG GTTAGTGGAT GTTTCGATCG AGAAATCTGC TACAATTAAA	4440
ACTACAAGTC ATTCAAGAAAT AAAAGAAATCC AAAGAAAGTAG TGAAAAGGA TCTTGAAAT	4500
AAAGAAACGA GTCACACATGA AAGTGTACA AATTTTTCAA CTAGTCAAGA TTTGACATCC	4560
TCAACTTCAC AAAGCTCTGA AACAGAGTGTAA AACAAATCGG AATCAGAACAA GTAGTAGAAA	4620
AGAAGGTTTT AGGGCCTTCT TTTCCCTATC AACTCTTTTC TATTTCTGT TATTGATGTT	4680
ATAATGGATA AATATGAATA ATCGGAGTGA GACTATGAAA TACAAACGGAA TTGTCCTTAA	4740
GGTGGGTACT TCTTCTCTGA CAAATGAGGA TGGAAAGTTTA TCACGTAGTA AGGTAAAGGA	4800
TATTACCCAG CAGTTGGCTA TGCTGCACGA GGCTGGTCAT GAGTTGATTT TGGTGTCTTC	4860
AGGTGCCATT GCGGCTGGTT TTGGAGCCTT AGGATTTAAA AAGCGTCCGA CTAAGATTCC	4920
TGATAAACAG GCTTCAGCAG CGGTAGGGCA GGGGCTTTTG TTGGAAGAAAT ATACAACCAA	4980
TCTTCTCTTG CGTCAAATCG TTTCTGCACA AATCTTGCTG ACCCAAGATG ACTTTGTGGA	5040
TAAGCGTCGT TATAAAAATG CCCATCAGGC TTTGTCGGTT TTGCTCAACC GTGGGGCAAT	5100
TCCTATCATC AATGAGAATG ATAGTGTGCGT TATTGATGAG CTCAAGGTTG GGGACAATGA	5160
CACTCTAAAGT GCTCAAGTAG CGGGCGATGGT CCAAGCAGAC CTTTTAGTTT TCTTGACAGA	5220
TGTGGACGGT CTCTATACTG GAAATCCTAA TTCAGATCCA AGAGCCAAAC GCTTGGAGAG	5280
AATCGAGACC ATCAATCGTG AGATTATTGA TATGGCTGGT GGAGGCTGGT CGTCAAACGG	5340
AACTGGGGGT ATGTTAACCA AAATCAAGGC TGCAACTATC GCGACGGAAT CAGGAGTTCC	5400
TGTTTATATC TGCTCATCCT TGAAATCAGA TTCCATGATT GAGGCGGCAG AGGAGACCGA	5460
GGATGGTTCT TACTTTGTTG CTCAAGAGAA GGGGCTTCGT ACCCAGAAAC AATGGCTTGC	5520
CTTCTATGCT CAGAGTCAGG GTTCTATTTG GGTTGATAAA GGGGCTGCGG AAGCTCTCTC	5580
TCAATATGGG AAGAGCTTTC TCTTATCTGG TATCGTTGAA GCAGAAGGAG TCTTTCTTA	5640
CGGTGATATC GTGACAGTAT TTGACAAGGA AAGTGGAAAA TCACTTGGAA AAGGACCCGT	5700

GCAATTGGA	GCATCTGCTT	TGGAGGATAT	GTTGCGTTCT	CAAAAAGCCA	AGGGTGTCTT	5760
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TTAGAGGTA	ACTATGGTA	GTAGACAAGA	ACAATTTGAA	CAGGTACAGG	CTGTTAAAAA	5880
ATCGATTAAC	ACAGCTAGTG	AAGAAGTGAA	AAACCAAGCC	TTGCTAGCCA	TGGCTGATCA	5940
CTTAGTGGCT	GCTACTGAGG	AAATTTAGC	GGCTAATGCC	CTCGATATGG	CAGCGGCTAA	6000
GGGGAAAATC	TCAGATGTGA	TGTTGGATCG	TCTTATTTG	GATGCAGATC	GTATAGAACG	6060
GATGGCAAGA	GGAAATCGTG	AAAGTGGTTGC	CTTACCGAT	CCAATCGGTG	AAGTTTTAGA	6120
AACAAGTCAG	CTTGAAAATG	GTTTGGTTAT	CACAAAAAA	CGTGTACCTA	TGGGTGTCT	6180
CGGTATTATC	TATGAAAGCC	GTCCAAATGT	GACGTCTGAT	CCGGCTGCTT	TGACTCTTAA	6240
GAGTGGAAAT	GCGGTTGTT	TTCGTAGTGG	TAAGGATGCC	TATCAAACAA	CCCATGCCAT	6300
TGTCACAGCC	TTGAAGAAGG	GCTTGGAGAC	GACTACTATT	CATCCAAATG	TGATTCAACT	6360
GGTGGAGGAT	ACTAGCCGTG	AAAGTAGTTA	TGCTATGATG	AAGGCCAAGG	GCTATCTAGA	6420
CTTCTCTCATT	CCTCGTGGAG	GAGCTGGCTT	GATCAATGCA	GTGGTTGAGA	ATGCGATTGT	6480
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278	
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280

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282

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284		
CGACTCGTCT	CATGGAGCGT	18120
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CCTTCAACT	CATGCTAGAA	
CACCTGAATT	CAGCCTATCA	18600
GATTCAAGCAC	CGAAAAGGTG	
GACGTCCACG	TAGTCTGCC	
ATGGAAGACC	AGCTCATTAT	18660
GACCCCTCCGT	TACTTGCAT	
ATTATCCCAC	TCAGCGTCTG	
CTGGCCTTTG	ATTTTGGCGT	18720
CGGTGTAGCT	ACGGTAAATG	
CCATCATCAC	TTGGGTGGAG	
GATAACACTTC	GTGCGTCAGG	18780
TAGCTTGAT	TTGGACCATT	
TAGAAGCCCC	GAGTGCTGCT	
GTGGCTATTG	ACGTGACCGA	18840
AAGTCCGATT	CAGCGTCAA	
ACAAAACCAA	AGCAAAATT	
ATTCTGGAA	AAAGAAACGA	18900
CACACCTAA	AAACTCAAAT	
TATGCTGGAT	TTGACGACAC	
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TTTTCTGACG	GACATACGCA	
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TGTTGACCTA	GGTTATTTAG	
GCATCTTGAA	ATTTCATGAG	19080
AATACTTTCA	TCCTGTCAA	
AAATTCCAAA	AATGCCGCC	
TGAGTGAGGA	TGATAAGCAG	19140
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GATACTGAATT	GAAATTGAAC	
ATTTAACCG	TAAATTCAAG	19200
ACCTTCCAAA	TCATGTCAGT	
CCCTTATCGT	AACCGCAGAA	
AACGTTTCGA	GTTACGGCG	19260
GAATTAATT	GTGCCATCAT	
CAATTATGAA	GTGAACCTAGA	
TTCCGAACAA	GTCTAATATA	19320
CTTTTGAGAG	AGGAAAATCC	
AGTTGTATAG	GCTAAAGGTT	
TTATCCAAAG	GTCTGAGACA	19380
ACGATTAGGC	ACGATGGAAA	
GAACCTTTAT	GTGGCTGATG	
ACGATCAGT	CATCTTCCTG	19440
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AGAAAAGTAGG	AATTGAAAAA	
GATGATTGAC	CAACTATCTA	19500
AGTATTACAG	TTGTAGGATA	
CTAACTGAAA	AGGATATTCC	
AAGTATTTA	TCTTTATATG	19560
AAAGTAATCC	TCTGTATTTT	
CAGCATTGTC	CACCGAGCC	
AAATTTGCA	ACTGTAAAAG	19620
AGGACATGCT	TTGCTACCT	
GAAGGTAAAG	CTAAGGCTGA	
TAAGTTTTT	GTTGGATTT	19680
GGAATGGATC	TGACCTTGTG	
GCTGTTATGG	ATTTTGTCTA	
TGCATATCCT	GATGAGGAGA	19740
CTGTTTTAT	TGGTTTGT	
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GAGAAAAGGG	ATTGGTACTC	19800
ATATTGTGAC	AGAACACTA	
GCTTATTTG	CTAAGAACCTT	
TCGAAAGGCA	CGTTGGCTT	19860

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GATGCGAGGT TAAGCAAGAA CTCTATACGG TTGTTATCGC TGAAACAGAGC CTAGAAGATT	19980
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TGGCGGCATT TATGACGATC GCTTTTAGT TAAACCCGTG CAAGCAGTCT TAGATAAGAT	20160
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GCCCCAAACCT AAAAAGAAAA AGCAAGGGTG ACGAAGTAA AAAAGAAGTC TGCTAAGGCC	20340
CTGTCCTTGC ACGGGTAAA TTTTATATAT AAAAAGAAGC TGGGACTAAA GAGCTCAGCT	20400
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TTATTCAATT TAGAGTTCT TATGCACAAT TGAGTCTGGA ACGAAAGTCT CCAGTTGCCA	20520
AGTATACAGT ACAATAAACC AACGATGTAA TAGCTGATGA CACAAAGCAC AGTGGGTAGG	20580
ACTTGCGAAG TCACCCCTTTT CTTTCAAAA TTTTACTAA ATCATTGATA TCAGTGTAGT	20640
CACGATTAAG TCCTTGAGCA ACTGGTAGGT TAGTCAGTA ACCTTGATAA GTACTCACAC	20700
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CGATATAAGG AAGAGTGACA TTGGTTAGGG CGATGGTTGA AGTGCAGCA ACCGCACCAAG	20820
GGATATTGGC AACGGCATAG TGAGAACAC CGTGTGTTTC ATAGACGGGT TCATCGTGCG	20880
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AGAGCCTGGA CGCATTTGTT TGACCATCTC ATCTGTCACC AATTCCGGTG CTTTGACACC	21000
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TGAATTAGAC ATAAGAGTTT GAATTTGACT TCCAAAGACT TCTTCTAGAA CTGAGAGACG	21120
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ATGTGTACCG ACGACACCAAC CACCGATGAT AGTTACTTTT CTTTTGGAA CACCTGGTAC	21240
ACCACCAAGT AGAACACCAAG AGCCACCAAGC TTGCTTAGTA AGGAAGTGAG CTCCGATTTG	21300
AACAGCCATA CGACCTGCAA CCTCACTCAT AGGAACGAGG AGCGGTAGTT GTCCTTGATT	21360
GTCACGAACA GTTTCAGTTG TTTTGCTGT TAACATAGCA TCTGCTAATT CTGGAGCAGC	21420
GGCCATGTGC AAGTAGGTGA AGAGAAGAAG ATCGTCGCGC AAGTAACCGT ATTCAGAACT	21480
TAAAGATTCT TTTACTTTCA CAACCAACTC TGCTGCCAA GCTTCACCAG CAGTAGCGAC	21540
AATCTCAGCT CCTTGCTTTT GATAGTCAGC ATCAGTAAAG CCAGAACCGA GACCAGCATT	21600

286

TGTTTCGATA	AGGACACGAT	GACCACGACT	AACTAACGCTA	TGAACACCTG	CAGGTGTGAG	21660
GGCGACACGG	TTTTCGTTAT	TTTTAATTTC	TTTGGGATT	CCGATTAACA	TTGAGATAAC	21720
CTACCTTTCA	ATTGACGGTC	TTGTTTGGT	TGTCACATTC	CAGTTCATAA	ATCAAAAATG	21780
TGACGGTTTC	ATTGTATATG	AAACCGCTTC	AAAATCAAG	AAAACATTGT	CATCCAAATT	21840
TTTTTATGCT	AGACTAGTGA	AAATCAAGCT	CTAATGGAGG	GAAAAGTATG	GAATCAATAT	21900
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CTTTGGATGA	TGCGGAacAA	TGTTTGACTA	TGCCTCGGAC	AAGGGTAATA	CACGTTACAC	22020
TTTTCCAACC	AATCAAAGCT	TGGAAGAAC	CAAGAATAAC	ATTGCTCAGT	TCTACTTGGC	22080
TAATCCCTTG	GGACGTTGGG	GAATAGAACT	AAAAGCAAT	GGTCAGTTA	TTGGAACCAT	22140
TGACTTGCAC	AAGATTGATT	CTGTTCTTAA	GAAGGCAGCT	ATTGGCTACA	TTATCAATAA	22200
AAAGTATTGG	AATCAAGGAT	TAACGACAGA	AGCCAATCGT	GCTGTGATTG	AGCTAGCTTT	22260
TGAGAAAGATA	GGGATGAATA	AGTGACTGC	CCTTCACGAT	AAGGCTAATC	CCGCGTCAGG	22320
AAAGGTCTAG	GAGAAATCAG	GCATGCGTTT	TTCCCATGCA	GAACCATATG	CTTGTATGGA	22380
CCAGCATGAA	AAAGGCCGAA	TCTTGACAAG	AGTTCAATTAT	GTCTTGACCA	AGGAAGACTA	22440
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AATAATCTAA	GAGAGGAGAA	AATATGGAAG	CAATTATCGA	GAAAATCAA	GAGTATAAAA	22560
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CTCCACAAAC	ACCTGTCAA	GAGACGAATT	TGCAGGCTGA	AGTTGCAGCT	GTTTCCAAGG	22680
ACTCATCGAC	CGAAAAGGAA	GTGAAGAAGG	AAGAAAAGGA	AGAACCCCTT	GAACAAGATC	22740
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GAGAAGAACG	AGTTAGTCAA	CAGACTGGTT	CGGGGACAGC	TTCTTCACAA	AGCAAGGAAA	22980
AGAAGGTCAA	TCTCAACAAG	GCCAGTCTGG	AAGAACTCAA	GCAGGTCAAG	GGACTGGGAG	23040
GAAAACGAGC	TCAGGACATT	ATTGACCATC	GTGAGGCAGAA	TGGCAAGTTC	AAGTCAGTAG	23100
ACGAGCTCAA	GAAGGTCTCT	GGCATTGGTG	GCAAAACAAT	AGAAAAGCTT	AAAGACTATG	23160
TTACAGTGGA	TTAAGAATTTC	CTCTATTCCC	CTAATTACCG	TGAGTTTTCT	ATTACTTTGG	23220
CTTTTATTAACG	CTATTTCTC	AGCATCTTAT	CTTGCTTGT	TGGGCTTTGT	TTTTCTGCTA	23280
GTCTGTCTCT	TTATCCAATT	TCCGTGGAAA	TCTGCTGGTA	AAGTTCTAAT	AATTGCGGA	23340
ATCTTTGGAT	TTTGGTTTGT	TTTCAAAAT	TGGCAACAGA	GTCAAGCGAG	TCAAAATCTG	23400

GC GGATTCTG TTGAAAGGGT ACGGATTTG CCTGATACTA TTAAGGTTAA TGGTGATAGT	23460
CTATCCTTTC GTGGCAAGTC TAACGGTCGT GCTTTCCAAG TCTATTATAA ACTCCAGTCC	23520
GAGGAGGAGA AAGAACCTT TCAAGCTTA ACTGACCTGC ATGAGATAGG ACTAGAAGGG	23580
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AAGACTCAGG GAATTACCA GACTCTCAAT ATCAAAACAA TCCAGTCACT TCAAAAGATT	23700
GGCAGTTGGG ATATAGGAGA AAACTTGTCC AGTTTACGTC GAAAGGCTGT GGTTTGGATT	23760
AAGACCCACT TTCCAGACCC TATGGCAAT TACATGACAG GACTCTTGCT GGGACATCTG	23820
GACACCGACT TTGAGGAGAT GAATGACCTT TATTCCAGTC TAGGAATTAT CCACCTCTTT	23880
GCCCTATCTG GCATGCGAGT AGGTTTTTC ATGAATGGAT TTAAGAAACT TCTCTTGCAG	23940
TTGGGCTTGA CCCAAGAAAA GTTGAATGG CTGACTTATC CCTTTTCCCT TATCTATGCG	24000
GGACTAACTG GATTTTCAGC ATCGGTTATT CGCAGTCCT TGCAAAAGCT ACTGGCTCAA	24060
CATGGGGTTA AGGGCTTGGA TAATTTGCC TTGACGGTGC TTGTCCTCTT TATTGTCATG	24120
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TTGGGCATAT TGCCATTCT ATCCTTCTAT TTTGCGGAAT TTCAACCTTG GTCTATCCTT	24300
TTGACCTTTG TCTTTTCCTT TCTTTTGAC TTGGTCTTCT TACCGCTCTT GTCTATCTTA	24360
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GGCATTATTC GCTTGGCTC GCAGGTGGCA AGGAGACCAC TTGTCTTGG TCAACCCAAC	24480
GCATGGCTTT TAATTTTATT GTTAATTCC TTGGCTTGG TCTATGATTT GAGGAAAAAC	24540
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ATGTAACCTGG GAAAACCATT CTCATAGATG TAGGTGGTAA GGCAGAATCT TATAAGAAAA	24720
TCAAAAATG GCAAGAAAAG ATGACGACCA GCAATGCCA CGAACCTTG ATTCCCTATC	24780
TCAAAAGTCG AGGAGTAGCT AAGATTGACC AGCTAATTTC GACTAACACG GACAAGGAGC	24840
ATGTTGGAGA TTTGTCAAGG ATGACCAAGG CTTTCCATGT AGGGGAGATT CTAGTATCAA	24900
AAGACAGTCT GAAACAGAAG GAATTGTGG CAGAACTACA GCGCACTCAA ACAAAGGTGC	24960
GTAGTATGAT AGTAGGGAG AACATTGCCA TTTTGGAGA TCAGTTAGAA GTTCTATCTC	25020
CAAGGAAAAT GGGAGATGGA GGACACGATG ATACCTAGT TCTGTATGGG AAATTCTTGG	25080
ATAAGCAATT TCTCTTCACG GGAAATTGG AGGAGAAAGG AGAGAAGGAC TTGCTGAAGC	25140

288

ACTATCCAGA CTTGAAAGTA AATGTTTGA AAGCTAGCCA ACATGGCAAT	AAAAAATCAT	25200
CAAGTCCAGC CTTCTAGAA AACTCAAAC CAGAGCTTAC TCTTATCTCA	GTTGGAAAGA	25260
GCAATCGAAT GAAACTCCCC CATCAGGAA CATTGACACG ACTGGAAGGT	ATCAATAGCA	25320
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TCGAAAGTGT TCGATAGGAA GGATAAAATGT TGTAGATTAG TGAAATAAC	AAAAATTG	25440
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CATTAAAAAT CAGCAAAAGT TGTTTATTA GTTAGTTTAT AATCTATTGG	TCTTCTTCAG	25560
TCCAGTGTAT CTGCTGTGAC AGTCACTAAA AGTTACAAAGT ATGATTGGAA	TACGGTTGG	25620
GAATATAGTA CCAACTATCA CGACCACATCG TATGCTTGA TTCCGTCATG	GTCTCGTTAT	25680
GACAGCTATT CTGAGTATAA AGTTGGCGGA GGCTGGAAC ACTGCTCGTTA	TGAGGTCATA	25740
AACTATTACA GCGGAGGCTA TTAATTCTTA AAGAGTGAGA AAAAGGAGGG	CTAGATATGT	25800
TGCAGCTTAC TCATGTGACC TTAAAAACGC GACAAGTCAT CTTGCAAGAT	GTGGATTCA	25860
CCTTTAAAAA GGGTAGGGTT TATGGTCTTC TTGCTATCAA TGGCTCTGGA	AAGACGACCC	25920
TGTTCCGTGC CATTAGCAAT TTAATTCCCA TAAGTAGTGG AAATATCGCA	GCCCCCTCCTT	25980
CTTTATTTTA TTATGAGAGT ATTGAATGGC TGGATGGAA CTTAAGTGGG	ATGGACTACC	26040
TTCGTCTTAT CAAAAACATC TGGAAGTCAG GTCTGAACCT GAGGGATGAA	ATCGCCTATT	26100
GGGAAATGTC TGACTATATC AGTCTTCCCA TTCGCAAGTA TTCCCTAGGC	ATGAAGCAAC	26160
GCTTGGTGTAT TGCCATGTAT TTCCCTCAGTC AGGCCAAATG CTGGCTCATG	GATGAGATTA	26220
CAAATGGCTT AGATGAGTAT TATCGACAGA AGTTTTTGA TAGGCTAGCA	CAAATCGATA	26280
GACAAGAACCA GCTGGTTCTT TTAAAGTTCCC ACTATAAGGA AGAGTTGGTT	GATGTCTGCG	26340
ATAGAGTAGT AACCAATTCA CAGGGGCAGA TAGAAGAGGT TTAGTTATG	AAAGATGTTA	26400
GTCTATTTTT ATTGAAAAAA GTTTCAAAA GCGCTTAAA CTGGATTGTC	TTAGCTTTAT	26460
TTGTATCTGT ACTCGGTGTT ACCTTTTATT TAAATAGTCA GACTGCAAAC	TCACACAGCT	26520
TGGAGAGCAG GTTGGAAAGT CGCATTGCA CCAACGAGAG GGCTATCAAT	GAAAATGAAG	26580
AGAAACTCTC CCAAATGTCT GATACCAGCT CGGAGGAATA CCAGTTGCT	AAAAATAATT	26640
TAGACGTGCA AAAAATCTT TTGACGCGAA AGACAGAAAT TCTGACTTTA	TTAAAAGAAG	26700
GGCGCTGGAA AGAAGCCTAC TATTGCGAGT GGCAAGATGA AGAGAAGAAT	TATGAATTG	26760
TATCAAATGA CCCGACTGCT AGCCCTGGCT TAAAATGGG GGTTGACCGC	GAACGGAAGA	26820
TTTACCAAGC CCTGTATCCC TTGAACATAA AAGCACATAC TTTGGAGTTT	CCGACCCACG	26880
GGATTGATCA GATTGCTGG ATTTAGAGG TTATCATCCC AAGTTGTTT	GTGGTTGCTA	26940

289

TTATTTTAT GCTAACACAA CTATTTGCAG AAAGATATCA AAATCATCTG GACACAGCTC	27000
ACTTATATCC TGTTCAAAA GTGACATTTG CAATATCCTC TCTTGGAGTT GGAGTGGGAT	27060
ATGTAACGTG GCTGTTTATC GGAATCTGTG GCTTTCTTT TCTAGTGGGA AGTCTGATAA	27120
GTGGTTTGG ACAGTTAGAT TATCCCTACC CAATTTATAG CTTAGTGAAT CAAGAAGTAA	27180
CTATTGGAA AATACAAGAT GTATTATTT CTGGCTTGCT CTTAGCTTC TTAGCCTTA	27240
TCGTCATTGTG GGAAGTTGTG TACTTGATTG CTTACTTTT CAAGCAAAA ATGCCTGTCC	27300
TCTTTCTTTC ACTCATTGGG ATTGTTGGCT TATTGTTGG TATCCAAACC ATTCAACCTC	27360
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GAAGATTACC TAAGCAGATT GATAATGTGC ATCTAAATTG GAGCATGGGA ATGGTCTTAC	27480
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CACAGAAAAA AGAATTTTTT AATAGATTCT AGCTTTCTA TAGGTAGGG AAATAAGTAA	27600
AAACTAACAT AGAGAGGGAA TCAACTTGAT TCTCTTTTG TGATTGAAA ACCAACCAA	27660
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GTATCATATA AAAGTTGAGA AAAGCAGAAG TGAGAGCTTC TCGCCTTGAT ACATTAAGTT	27780
GCCTGGCCCT ACGGATGAAA AGTTTCAAG AAACGCTATC ATAACGTGCG GGCTGTATA	27840
TTTACAAGTC CGCTATTGTT TTTCTCTAAT AAAACAAAAG AGGTGAAAAC CATAGCAAAG	27900
CAAGACTTAT TCGTCAATCA TCAAGATTCTG GTACGTGAAG TTGCGTTGAT TGGTCTTGAA	27960
GGAGAACAGC TAGGTATCAA GCCACTCAGT GAAGCGCAAG CTTGGCTGA TAACGCTAAT	28020
GTTGACCTAG TATTGATTCA ACCCCAAGCC AAACCGCCTG TTGCAAAAT TATGGACTAC	28080
GGTAAGTCA AATTGAGTA CCAGAAGAAC CAAAAGAAC AACGTAAAAA ACAAGCGTT	28140
GTTACTGTGA AAGAAGTTCG TCTAAGTCCG G	28171

(2) INFORMATION FOR SEQ ID NO: 23:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 7147 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 23:

CCGCTCAACT TTTGCAATCA AGGCTAAGTA GACAGCAGCA AATTCATAT TGTATAATT	60
CTGACTCATA CTTCTCTCTT TCTATGTGTA CTAGTATAAA TAAGAAAAAG AAGGCCGTCA	120

290	
AGCCTCTTT TGATTTATTC TTCTGCTTCA TCTTCTGTAATTGACTATT GTACAAGTCA	180
GCGTAGAAC CACCTGCGC CATCAGTTCC TCATAGTTGC CTTGCTCGAT GATATTTCCA	240
TCTTTCATGA CCAAGATCAA GTCTGCATTG CGGATGGTTG ACAAGCGGTG GGCAATGACA	300
AAGGATGTGC GTCCCTCCAT CAAACGGTCC ATGGCTTTTG GGATCAATTCTCTGTCCGT	360
GTGTCAACAG AAGAAGTCGC CTCATCCAAA ATCAAAAGCG GTGCATCCTT AAGAAGGGCA	420
CGAGCAATAG TCAATAGTTG TTTTGTCCTT ACAGACAAGG TCACGGTGTCA ATCCAAGATG	480
GTATCATAGC CATCTGGCAA GGTCTATAATA AAGTGGTGAA TTCCCACAGC CTTACTAGCT	540
TCCATCATTC GTTCATCACT AATCCCTATT TGATTATAGA TGAGATTGTC TCGAATAGTT	600
CCTTCAAAGA GCCAGGTATC CTGCAAGACC ATTGAAAAGG CATCATGCAC TTCTGAACGC	660
GTCATAGCCT TGGTATCCAC ACCATCAATG CGAACTACTTC CCTTATCAAT CTCATAGAAT	720
TTCATCAAAA GATTGACAAT GGTGTCCTTA CCAGCCCCAG TCGGGCCAAC AATGGCAACC	780
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CCAGAACCTG TTGACCCAAAT AAAGGCCACT GTTTGACCAAG TTTCTGCTTT AAAGCTAACAA	960
TGTTCAATAA CTGCTCCGA ATTTGCCGCA TAGCGgAAGG TCACATCCTT AAACTCGACC	1020
TGACCTTTGA AGTTTCATC AGTCAGCTGC ACTTGAAACAG GGTGTTGGAT AGAAGAATGC	1080
AAATCTAAA CTTGATTAAT CCGCTTAGCA GAGACCATAG TTGGGGAAAG AACGATGAAG	1140
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ATCCAGTAAA TCGCCACACT CAAACCACTT GAAATCCCCA TCATGATAGG ATTCAAAATA	1320
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TTTTCATTTT GATAATCCTC TGCATTGTCAG GCACGAACGA CACGAATACC TGTTAAACTC	1440
TCACGAGTGA TACTGTTCAAG TTTATCTGTC AGCCCTGAA TCAAGGACTG TTTTGAAAG	1500
GCTAGCGTCA TCAAAACGGT CGTCATCAGG ACGTTGATAA TCACGCCAC AAGTACGGCC	1560
CAGAGCCAGT ATTCTGAATG ACCTAAAATC TTCCAATAG CCCAGATAGC CATAATTGAA	1620
CCACCGGTTA CCACCTGCAA GCCCATAGTA ATCAACATTG GAACTTGAGT AATGTCATG	1680
GTAGTACCGG TCAAGAGGCT AGGAATTGAA AATTCTTAA TCTCTGTCG CGAGTAATCC	1740
AAAACCTCGGT TAAAAATATC ACTTCTCAGC CTACTAGTAT AAGAAGCCGC CACTCGGGAT	1800
GCAAAAAATC CAACTGCAAC TACGGACAAG AAGGCAAGAA AGGACATTCC CATCATCATG	1860
CTTGCCGACT GCCACAACTC ATCTAAATTAA GTTTCTTGAC TACCTAGCAA ATCCGTAATT	1920

TTCGAGATAT AGGTCGGCAC TTCCAATCT AGATAGACCG AAAAGCAAGT AAAGAGAATG	1980
GCTAGTAAAA TCATCCCCCA TTCTTTCTA CTAATTCTTT TGGCTAATTT CTTTATTCTC	2040
TCCTCCCTATT CCCTTGATAT TTTGCCTGTA GTTGACCGAG AACCTTCTCA AAAATCAGTA	2100
ATTCATCTTC ATCAATGTCT TCCATCAACT GCTTGTCTAT GCGTTCAAAA AAAGCCTTAA	2160
CCTGTGCGAT CTGAGAACGT GCTTGTCCG TCAGACGAAC AAACCTAGCC CGCTTATCAA	2220
CAGGACTCGC CTCCAATTCC ACCAAACCAT TTTGCACTAT ACGCTTAACC AGATTACTAG	2280
CAACAGGCTT GGTAAATATTG AGTTCTGCT CGATATCTTT AATCAAGACC AAGTCTTGGT	2340
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AGACTAACAT CGGTTTATCC ATAATCTCCC CCTCTAAAT AAAATAGTT CTCTGGAGAA	2520
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AGACTGCGTC CGCATCTTCA CCTCCAAATA CAATAACACC TTTGTTGTC AAGAGGTAAG	2940
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TCGCTTTATC GATAATGTGT TTTGGCACTT GGGCTTGTGTT AGCACGGTCG ATAACGAATT	3180
TCAAAGCTGA GTTTGATTCT GGATCTGGAT CACCTTTTTT AGCTGCTACA TAGATTCTA	3240
CACCAAATTT TGCATATACT TTAGAGTTAG CTCCATCTTT AGCCGTTTC TTGGCTACGA	3300
TATTGGCCA TTTACGTCCC ATTAGGAATC TCCTTTTTTC ACATTTTAAT CTTTCTTATT	3360
ATAACACAAG TTTTTTGAT TTTCACTAGA GGAAATGGAT TTTATTAGCA AATCAAGCTA	3420
GGATAGCACT TTACCTGCTA AGATGGTCTT GCCTTTCTAT CTTTATCAAC AGGCACTCAT	3480
CCACATTCAA AAAACAAACT AGACCATTAT CTGCAAATAG AAAGTTTCAG CCAAGTTGA	3540
CAAAGTCAGC TCAAATTACT GTTGAGTT TGTAGATATA AGCGACAAAA ACAATCATA	3600
TGCACCTTTT GTTGACAGTC TACTCCAGAC ATATCATAGT TCAAGTAAAT ACTTTGAAAT	3660

292

TCAACAGTTC TTATAGCGC TATTGTATTC TAAGAAATCA ATAGAAGAGT TTCTAAGCAA	3720
ACCTCTAATA CTCAAAAAA ATCAAAGAGC AACTAGAAA GCTAGCCTCA GGTTGCTCAA	3780
AAACACTGTT TGAGGTTGCG GATGGGGCTG ACATGGTTG AAGAGATTT CGAAGAGTAT	3840
AATTACGTG TTCCCAAGAT GGAGAAGTTA GACTAGTACA CTGGCACTTC TAAAACATTG	3900
CTAGCAATTG ATTTGTTCAT ATTTAATTTC ATTTTTCCA TAAATGGTA TTAGATATAA	3960
ACAGCAAAAT ATTTCCGATA CGTGTGTTTC TTGAATTCC AATCATCTAA ACAAGTAAA	4020
GGATAATCAA TCCCCTGTAT ATCAAGGAAT TGGCTACCCCT TTTTACTTTT TTACACATTG	4080
TGTTTGATAG ATTCAATTAA ACATCACGAG CATACTCCAA TGGAAATCGC TAGGCAAGAG	4140
ATAAACATTG AGATATCCGC AGAGAGATCA TCGCCTCTTT TTGTCGCAAG CATTCTCCTC	4200
TCCTAGTCAT TTTCTACCTT ATCTTCTACC TGAGGATAGA GAGTTGTTCC CCAAATAGAA	4260
ATCGTCCGCT TACGCACTAG TGGCAAATCG GTTTTTCAT AAACCGTACG CCACCATTC	4320
CAGGCAAGGCC CGGTACACTC TCTAATTTC ACAGAGAGAT TACGAACATT CCCTTTAAA	4380
GGAATACTAG TGGTAAAGTG AGCCGTTAAA TCCTGCCAT TTCTGTCCCA AGCCTTAGGA	4440
GTCAAGACTT CCTTACCTTG ATGATCATAG GATAATTCTAT TCCAAGTAAT ATAATATTGG	4500
GCAACATAGG CACCACTATG ATCCAGCAGT AAATCTCCGT TTCTGTAAGC TGTAACCTTA	4560
GTCTCAACAT AGTCTGTACT ATTTGAAAG GTCGCACTA CATTGTCAG TAAAAAAGAA	4620
GTTGTATAGG AAATCGGCAA GCCTGGATGA TCTGCTGTAAGCGACTGCC TTCTTGAAATC	4680
AAAGTCCTCTA CCATATCCAC CTTGCCTGTT ACAACTCGGG CACCCGAACACT TGGGTCGCC	4740
CCTAAAATAA CCGCCTTCAC TTCTGTATTG TCCAAAATCT GTTTCACACTC TGTCTGAGGA	4800
GCTACCTTGA CTCCCTTTAT CAAAGCTTCA AAAGCAGCCT CTACTTCATC ACTCTTAC	4860
GTGGTTCCA ACTTGAGATA GACTTGGCGC CCATAAGCAA CACTCGAAAT ATAGACCAA	4920
GGACGCTCTG CAGAAATTCC TCTCTGTTT AAATCCTCTA CCGTTACAGT ATCTTGAAAC	4980
ACATCTCCTG GATTTTAAAC AGCATCTACG CTGACTGTAT AATAATCTG CTTAAAATTA	5040
ACAATCTGAA TCTGTTTC GCCTGAATGG ACAGAGTTAA AATCAATATC AAGAGAATT	5100
CCTGTCTTT CAAAGTCAGA ACCAAACTTG ACCTTGAGTT GTTCCATGCT GTGAGCCGTG	5160
ATTTTTCTAT ACTGCATTCT AGCTGGGACA TTATTGACCT GACCATAATC TTGATGCCAC	5220
TTAGCCAACA AATCGTTAC CGCTCCCCGA ACACCTGAAT TGCTGGGTC TTCCACTTGG	5280
AGAAAGCTAT CGCTACTTGC CAAACCAGGC AAATCAATAC TATAAGTCAT CGGAGCACGA	5340
TCGACCGCAA GAAGAGTGGG ATTATTCTCT AACAAAGGTCT CATCCACTAC GAGAAGTGCT	5400
CCAGGATAGA GGCGACTGTC GTTGGTAGCT GTTACAGAAA TATCACTTGT ATTTGTCGAC	5460

293

AAGCTCCGCT	TCTTCTTTG	GATAACAACA	AACTCATCGG	GTAGCTGATT	ACCCTCTTGT	5520
ATGAAACGAT	TTTCAATACT	TTCTCCCTGA	TGGGTCAAGA	GTTTCTTTTT	ATCGTAATTG	5580
ATAGCTAGTA	TAAAGTCATT	TACTGCTTGA	TTGCCATCT	TCTACCTCT	AATAAGTTCC	5640
TGGATTGAGT	TGCATAAACT	CAGACTTGTT	CAGCGAAATC	AGCCGTGGTT	GGACTAAGTA	5700
ATCCAAAATT	TCCTCGTACA	ATTCTTCTGA	GACATTGCGT	CGCCGTCTGG	CTAAATAAGA	5760
AGTCGGAATG	ACCGTATTAT	CCAACATAAA	TACCTTATCT	AAGTCAATCA	AGGTTGGTCT	5820
TGTAAAAGGA	TTACGAGCTA	GATCCGGCTC	TTCTATCATA	AAGTTCTTGA	CCAAACGTCT	5880
GGTCAAGAGA	GCTGGTTGA	AGGCTCTGATT	TTAACCAAC	TCTTGTGTTT	TAGTCATGCT	5940
GTTGTCATAA	CAGATATACA	TATGATTCTT	CACAGCCAAA	TCGCTACTAA	TAGTCGGAAA	6000
AGGCAAATAA	AGAGCTACAA	CATCTCCTCT	CTTAATCAAG	CAAGAGCACC	CCCTTTCTC	6060
CTAATGTAAC	ATAGACAGGA	TTGACCAAGT	CTTCTGATTG	ACTCAGAATT	TCCAAAGTTT	6120
GAGTTGGCG	CGCTGTCAAT	TTAGTAGCAT	CTTGTCTCTT	CAATACAAAA	TGCTTGTGCG	6180
CAATAACCTT	GACAATATAA	TCTTCTCCA	AAGCTGACTG	GTAAATCCAC	ATCAGATGTT	6240
GTCTGTCTG	AGAACTCAAG	AGAGAAGGAT	TTTCAAGCCT	CCCGATAGTC	TGATAAAAAT	6300
CAAAAACAGG	AGCTAACTCC	TGCCAATCTG	ATTGGCTAGT	TGTCAAGGCT	AGAAAAAGGG	6360
CTTTGCGAGC	TGATACTCT	TGGTTAGCCT	TGAGAGTTAC	TTTCCCCTCC	AACTTTTTA	6420
GAANTCCCCA	AACTCCAGAA	AGCAAAATTTT	TCTCTAACTG	CGAGAAATAA	AAACCTTTG	6480
TTCCCAGACA	TAAGTCTTTC	ATGTCGCTTT	CTCTAGCAA	TAAGAGCTCA	AACATTTGAT	6540
AGTAAAAGAA	AAATATCTGG	CACTGGGTG	CGCTCATCTT	TTCTTATCG	GCTTCTTTT	6600
TTAACCCAGAG	CAAGGGCGAC	AGGTAGCTGG	ATTGAGACAT	TTCTCTTAC	TCCTACTCTT	6660
TTTTAACTGG	AGCATCTGCA	CTAGCTGCCA	CTTCTTTGA	CTGGATACCT	TCCCACGTG	6720
TAATCTCCTC	TGAGATAAGA	CCTTCGCATG	TCTTGACAAA	TAGGGCAAAA	GCCTTGGTCT	6780
TTCCCTGCATA	TTTCTCCGTT	TGGCATTGAT	AGAGGAATT	TTCTTCTCC	AGGAGTTGCG	6840
CAGTTTTTG	GTAAAGAAATC	CAATTTCCT	TTGCATTATA	CAAATTGATA	ATCCCCTCAC	6900
ACAGCAAGCC	GAGACTGGAT	AAGGCAACCG	AAATCAAACG	GTAGCGATCA	CCTGGCATAG	6960
GAATAGCACA	AAAGACAGCT	ATGAGGAAAC	CTGCCACGAT	TTCTGTTATT	TTAATACCT	7020
TATAGCCCT	ACGATGTTGA	ACGCTTTCT	TTAAAAAAATG	AGCTATCTGT	ACGTCTAATC	7080
GCTCTGTCAG	GTACATTTCT	TCTGGCGTCA	TATTCGTAAC	TCCTTCATT	TACTTTGATA	7140
ATCAGGG						7147

294

(2) INFORMATION FOR SEQ ID NO: 24:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 755 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 24:

CCGCATGGGA TTGGTGTCT TTTGGCAAT CTCTTGACC	AAACTGGAAA CATGTTTAT	60
GCGCCTGCCT TTACTGCCCT TGTCGGCGGT	ACGTCTATAT GATCCTAGTC	120
CGCGCTTGG AGCCATTACC	ACTATCGGCC TTGTCATTGC	180
AACACGGTGC	TGGTCCCTTC CTTCTGGAA	240
TTATCTGTGG	CCTCTTAGCA GATGGAGTAG	300
CTCATTTAGG	AAATACAAG GACAAAACAA	360
AGAACATTCT	TTCTTTCTATT ATTTTCGCCT	420
TTAGTACAAC	AGGACCAATC TTGCTTATGT	480
GGATTGCGCC	CAAAGCCTAT ATGGCTACTC	540
TTCTGGCAAG	AGGAAAATCC CAAGAATATA	600
TCGACCGTAT	CATGGTCGCT CAAACCCCTG	660
GAACGTGCCT	TCTATTTATC GCAAGTATTG	720
TCATCGGAGC	CCTAGTGGGT GCCTTGATTG	780
GACAAGCCTT	GAGTAAAAAA TTTGCCAGA	840
AAATCTGATC	AGTTAAAAAG AGCCACGCGG	900
CTCTTTTTA	TTTATGGCTC	960
AAATTCTTAG	TCAAGAAATC	1020
TCCAAGAAT	TCCCAAGAAT	1080
AGATAATCAA	TGGATAATA	1140
ATGGTTGCCA	AGATGGTCAC	1200
ATCGTGATTG	TAGCGGTTAA	1260
ATCCATAAGC	GATGGCTACG	1320
TTACCGATAC	CACCAAGCTCC	1380
AACCGCACCG	GCCATAGCTG	1440
TTtccAAACA	AGGGAAAtCAA	1500
GGTcACAGTC	GTCAC	1560

(2) INFORMATION FOR SEQ ID NO: 25:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 3010 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 25:

TTCAATTGGT ATCTCAATCA	ACGGTCTTCA CATGGTTCA	ACTGGTTGA CTCTTGAAAA	60
AGCGAAAGCT	GCTGGTTACA	ACGCAACTGA	120
AACAGGCTTT	AACGATCTTC	AAAAACCAGA	180
ATTCATGAAA	CATGACAACC	ATGAAGTAGC	240
AATTAAGATT	GTCTTTGACA	AAGATAGCCG	300
TGAAATTCTT	GGTGCCAAA	TGGTTTCACA	360
TGATATTGCA	ATTAGCATGG	GAATCCACAT	420
GTTCTCACTT	GCTATCCAAG	AGCATGTGAC	480
AATTGATAAA	TTGGCATTGA	CAGACCTCTT	540

295

CTTCTTGCCA CACTTCAACA AACCATAACAA CTACATCACA ATGGCTGCCCTT	360
AAATTAACCA TGAATGAGCT ATCTGGCCTT AAGTTAAGGT CAGATAGTTT TTAGCTAATT	420
TGTCCCCATA CAATTATAGT TTTTTTATCT TGTGCTTCAT TCTGTTCTGA CTTAAAATGA	480
AAAGGTAGCT ACCAATACAA ATGATGAGGA TAAAACAAAT GACTGAAAAT CGTTATGAAC	540
TAAATTAACCA CTTGGCACAG ATGCTCAAGG GTGGTGTAT TATGGATGTG CAGAATCCTG	600
AACAGGCTCG TATCGCAGAA GCTGCTGGTG CGGCAGCTGT GATGGCCTTG GAACGAATTG	660
CGGCTGATAT TCGTCAGCT GGAGGAGTTT CCCGCATGAG CGACCCAAAG ATGATTAAGG	720
AAATCCAAGA AGCGGTTAGT ATTCAGTAA TGGCTAAGGT CAGAACATCGGG CATTGGTTG	780
AAGCTCAGAT TTTAGAGGCT ATTGAATTG ATTATATCGA CGAGAGTGA GTTCTATCTC	840
CAGCTGATGA CCGTTCCAT GTGGACAAGA AAGAATTCCA AGTTCCCTTT GTCTGTGGTG	900
CTAAGGATTT GGGTGAAGCC TTGCGTCGTA TCGCTGAAGG TGCTTCCATG ATTCTGTACCA	960
AAGGAGAACC AGGGACAGGG GATATCGTCC AAGCTGTTG TCATATGCGT ATGATGAATC	1020
AGGAAATTG CCGCATTCAA AACTTACGTG AGGACGAGCT TTATGTTGCT GCCAAGGATT	1080
TGCAAGTCCC TGTAGAATTG GTCCAATATG TTCATGAACA TGGAAAATTG CCAGTTGTA	1140
ATTTCGCTGC TGGAGGTGTT GCAACGCCAG CAGATGCTGC GTTAATGATG CAATTAGGG	1200
CAGAGGGGGT CTTTGTGCGT TCAAGTATTT TCAAGTCAGG AGATCCTGTT AAACCGAGCGA	1260
GTGCCATTCT TAAGGCTGTC ACTAACTTCC GTCATCCTCA AATCCTAGCT CAAATCTCTG	1320
AAGATTAGG AGAACCCATG GTGGTATTAA ATGAAAATGA AATCCAAATT CTCATGGCTG	1380
AACGAGGAAA ATAGATGAAA ATCGGAATAT TGGCCTGCA AGGGGCTTT GCAGAACATG	1440
CAAAAGTGCCT AGATCAATTAA GGTGTCGAGA GTGTAGAACT CAGAAATCTA GATGATTTTC	1500
ACCAAGATCA GAGTGACTTG TCGGGTTTGAA TTTTGCCCTGG TGGTGAGTCT ACAACCATGG	1560
GCAAGCTCTT ACGTGACCAAG AACATGCTAC TTCCCACCCAG AGAACCCATT CTATCTGGCT	1620
TACCAAGTGTGTT TGGGACCTGT GCGGGCTTAA TTTTGCTGGC TAAGGAAATC ACTTCTCAGA	1680
AAGAGAGTCA TCTAGGAACATGATGAGCG TGTTGAGCG AGATGCTTAT GGGGCCAAT	1740
TAGGAAGTTT CTACACGGAA GCAGAAATGTA AGGGAGTTGG CAAGATTCCA ATGACCTTTA	1800
TCCGTGGTCC GATTATCAGT AGTGGTGGTG AGGGTGTAGA AATTTAGCA ACAGTGAACA	1860
ATCAAATTGTTG TGCAGCCCAA GAAAAAAATA TGTTGGTAAG TTCTTTCTAT CCAGAATTGA	1920
CTGATGATGT GCGCTTGCAC CAGTACTTTA TCAATATGAG TAAAGAAAAA AGTTGAGATT	1980
GAATTCTCA ACTTTTTTAC ATGTAATAAA CAATAGCGAT GTATTGAAGT GCGGACGCGAG	2040

296	
CTAGGATAAA GAGATGCCAA ATCATGTGGA AATAAGGTTT TTTCTTGGCA TAAAATCCAG	2100
CTCCAACGT ATAACAGAGT CCCCAGTTA CCATGAGACT CCAGAAAACG GGTGTCGTTT	2160
GAATGATAAT GCCAGGAATG ATAGCCAGAA CCAACCAGCC CATAATCAGG TAAAGAGCAA	2220
GGCTAAATTT CTCATTGACC TTTTAGCAA AGATTTATA GAGAATACCA AAGATGGTCG	2280
TTCCCCATTG GATGACAATA ATCAGATAGC CAAACCAGTT ATTCATCAAG GTCAAGACAA	2340
CGGGCGTGTG TGAGCCGGCA TGAGCAACGT AAATCATAGA ATGGTCAATG ATTCGCAAAA	2400
CATATTTGTG GGTGAAACCA TAGGCCATAG AGTGATAAT GGTTGGATGAT AGGAACATGA	2460
GAAAGAGACT GATGACGAAA ATGGAAACGC CGATAGAGGA TAAAATCCG TGTGCTTCAT	2520
AACTATAGAT GGATGAAATA GGCAGCAAGA TAAGCATGAT GACTGCACCC ACAGCATGGG	2580
TCACGCTATT AGCAATCTCC TCTCCAAAAC TGAGTTGTTT GCTGAGTTTA AGACTAGTGT	2640
TCATGGATT ACCTCCTCTT GAGTATGATC GATTAAGTCT AGAGTTTGAT GATAGAGTTT	2700
AACGGTTGG CAGCTGGTTT GGATAATAGG GTTAGCTGGG TCAATTCCCTT GGTCATGTA	2760
GTCCACAAAA GCATCGTAGA GTTGGTCTGA ACTTGCTTGA GTTTGTAGAG TATTAAGTGT	2820
CTGGGCTATT TCTTGAATAG AAAATACAGA CTTGAGGGTT GTGATAGCAA TCAAACGGGC	2880
AATCTGTTGG CGTTGGTATT TTTTTTGTC AGGCTTTGTC AGGTAACCAT TTTTCACATA	2940
ATTGTTGACC ATAGATGCTG TTAGGCCCTT GTCTTTATTA GGAGAGATAG GGGCGCAGAC	3000
CTGATTGACA	3010

(2) INFORMATION FOR SEQ ID NO: 26:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 15213 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 26:

CATAAACCGG TGCAAATAAC TTAATAGTGA AGTAGCCATT TCTTCGTAT TTACCTGAGG	60
CATATTCCCT AGACGAAAGA ATATTATTAT CAATCAAATC ATTGAATGAA CGTAGTCTTT	120
CAACTCTTC TACTGTTAGA TTTCTGACAA CATTGTTGC ATAGACCTTA TTTCCATCAG	180
GATCAGGATG GTACTCATTT GAACTTTTC TAAGAAGTTG TTGTTTTGAA TTGATATCCA	240
ATTTAAGAAT TGAATTTCTC TCGAGATATT CCAACATATA AACAAACGTCA AACATGTTGT	300
GGACATATTG CTTCAAATCA TCTGCATTAT TAAATCTTGT AGTTGGATCA AGTACTTGT	360
ATCGTCGACT TTCTGTACTA TCAGATTTTG AATGTTCAA GATGGAGTTG ATGGTAATGG	420

TCGCATCATC	TGGATGGTCT	GGTGCTTGTA	ATAATCCTTT	AGCAAAGAAC	TCTGGTCCCA	480
AGCCACTTCT	TCGACCATAT	CCTCCAAGAT	AAATGTCCTG	ATCTGAGTCA	TGTGTCATCT	540
CATCGGTATA	AGTAATAGCT	CCATCCTTAT	CCAACATTG	ATAACCCATA	TAATAAACTG	600
CATCACCTGT	AGCATAAGCA	CCGTGTTGAT	TATGCCAAC	TTTATTTCCA	ACAGGTCCAA	660
AGAAAATGTTG	CATTGCAGGA	TTTGGATTAT	CAAATCTGC	CACTTCTGTA	GCTTTCCCTA	720
CGGTATTATC	ATCGCCAAAT	TTATAAGCAT	CGTAAAGCAA	AATATTTCTA	AAAAGTTTT	780
CACGTGCATT	GTCGTCTAAA	ATACGATACC	ATAATCGTA	GTGATCTCGC	TGACGTTGG	840
CTGTTTCACG	CGCATTTCCT	TCAACAAAAT	CATTGAGACC	CTTGGCCGCT	TTATGGTCAC	900
TACTGCGGTA	GCGATCATAA	GCTCCAAATC	CTAGACTAGA	CATGGTCGAG	ATGACAAATA	960
CGGATCTCTC	TGGCAAGGTC	AGGAGAGGCA	AGACCATATT	GCGGTATTTC	CATGTGGCAC	1020
TCGTGATACG	ATCATAAAACA	CCGATAGAAC	ACTTGGTGCC	AGCTAACCC	TGCTTCGTTT	1080
TCACCTCTTC	GATAGTGGAT	TTTCTTCGA	CAATGTAAGC	CTTAGTCTCT	GATTTAAACC	1140
AGTCATTATT	GCTTGTATT	GGTAAAAAGA	CTTTTCGGTA	ATGTTCCAGC	GTGCTAAACA	1200
AATCTGTCGT	TCCATGTTGA	CTGGCAAGAC	TGATACCATA	AGTATCGACA	TTATTCTTAG	1260
CTAGAAAGATT	GTTAAAGCCA	GATTTACCCA	ACTCAATCAG	AGTATCTAAT	GGTGAAGCAT	1320
TCCCCTTACC	AAAGAAGTC	AAATGGTACA	GAACTAGGGC	TTTGACATT	ACCTGACCAT	1380
AGCTAAAGTT	ATACCAACCT	TCCACATAGC	TCAAGCCAAG	TAGCAAGGCT	TCCTTGTG	1440
GTGGATT	ATCTACAAGA	TAACCTTCAG	TGACGGGGTT	AGCACTAGCC	AGTCCAGCAT	1500
CCGCTGACAA	GAGTTTTTC	AAACTGTCTT	CCAGTTGTTG	TTTGTGTTTG	GCGAACTGGT	1560
CTTCTAGATA	GAGCTCAGTT	TGCTTGACGT	TTGGAGAAAT	ACCCAGCGTC	TTTCTGATGG	1620
CTTCTGAATG	ATAGTCAACC	TTTGTAAGT	CAGGTAAGAC	TTGCTTGATG	ATAGAGGTTT	1680
GGTCATACAG	GAATTGGTT	GGCGTATAGA	GAAGTCCAGT	ATTGCCAGA	CTATATTCTG	1740
CTAATTGGC	GAAATCATT	TGGTATTTGA	GATCCAGCTT	CTCAGATAAA	TCATCCTTGT	1800
AGTGAAGCAA	GAGTTGTT	GCAGTCTGTT	TGTTAGAAC	AATGTCCTG	ATGACTTG	1860
TGTCCTTCAT	CATGACTGCT	GACAAGAGTT	CTTTTGATA	TAAAAGACTG	TTCTCATTGA	1920
CCAGGTTTCC	GTATTTGACG	ATGGTTGCCT	TGTTGTAGAA	AGGTAGCAAT	TTTCAATGT	1980
TTTTATAAGT	CAAGTTGCGC	TTAGCTTGAT	AATAGGCCAC	CTTAGAAAAA	TCACTGTCTT	2040
TTTTGCCACT	TGTTGAAAGT	GGCTCCACTG	TTGGTAAAT	GAGAGGATTG	ATTTCTGCTT	2100
TTTTGCTTGC	AATTGAGAA	GCATCTAGCA	TTGTTCCCTCT	TTCTTCAAAG	GATTCCCTGC	2160

298	
TGACGACCTC ATCCTTGACC AAGGTGACAT TGTAGACTCT GTTGGCCTTG CTGCTGAATG	2220
TGTCCTTTAC CTTCATTTCG TTATAGTGGT AACAGTGAT GGCATTTCCG TTGGTTACAT	2280
TAACATCGCT GAGAACATTG GTCAAACCTTC CAGCATGCCT AACATCACCA GAAGTTCGAT	2340
CCCACAAATT GCCTGCCACT CCAGCGACTC TACCAAAGTG CTTGACATTC TTGATATCAC	2400
CTTCAGCATA GCTATCTTGG ATCTGTGCAT CTCGGTCTAC TAGGCCTGCA AGTCCACCCA	2460
CAGTCGTGATC TGAAGTATTG GTGTTAGATG AAATGGCTAC TGTCGCTTT GACTTAGTAA	2520
GTAAAGCCTT GTCACCTGTC AAATGACCGA CCATACCACC GATATTGTAG GCAGCAGTCG	2580
TTTCATAAGT GTTGATAATT CTTCCCTTGA AACTGCTCTC TGTGATGCTT GATTGCTCAG	2640
CCTTAGCCAG CAAACCACCG ATACCACGTT CACCAGCCAG AACACCACCG ACAGTGAACCT	2700
GCTTAATTTT TGTTGTTATTC TGAGCTTCAT TTGCCAGTGA ACCGATATCA TCTTTCCCTG	2760
AAATAGCAAC ATTTTTTAACT CTCAGTTTTT CTACTGTAGC ACCACTCAAG TTTTCAAACA	2820
GAGGTTTTTT CAAATTATAG ATAGCATAAT TCTTGCCATC TTTTCACCG ATTAACGAC	2880
CAGTAAAGGT GTCCCTGATA TAGGATCTTT CATCAGGACC AAGCTCCACT TCGTTAGCAT	2940
TCAGGGCTGGC CGCTAAATGA TAGGTTCCAG AGGGATTTG GTTGTAGCT TTGACCAGAT	3000
TAATGAAAGGAG AGTAAAGTTT GTTGTTCCTT CTGTTCCCTT CTTAGCTAGA TAGAAGGTAA	3060
AATTATCTTT ATATCTGCTT TCTATCTCCT GCTGAAGCTT CTCTACTTTT GCTGTGATTT	3120
TATAAAGGAT TTTATCATT TTTCTTCCT CTGATATTGA TGCTACTGGT AGGTATACAT	3180
CTTTGAATGA AGAAGATTTTC ACTTTAACAA AGTAGCTATT TGGATTGCTT GGAACCTGCT	3240
CTAACGAAAT GTGTTGTTA TAAGTACCAT TTGACAAACT GTATAACTCT AGGTGGAAA	3300
CATTTCTTAA TTCAAGTGTGTT TTCTCTGGTT CTTCTACCTT TTTATCAGGG TCTAGTTCAT	3360
TTTCTGTTT AATTCTTCG TTTCCATTG AATTGGATGT GTTTGATTG GTTGAAACAT	3420
CCTCAGTTGA ATTTCCGTTT GATGGTTCTG GTTCTGTTG TCCATTCTCT GATGTTGTAT	3480
TACCTGAATT TTCTGGTTTT GTTGCAGTTC CGTTTTTTTC TGTTGATTT GATTCTCAA	3540
CTGGTGGTTT TGAATCACTA GGTTTATTGG ATACTCTCC AGTATTTCTG TTAGCTATT	3600
TCCCAGAGTT TGTTGTGTT TCTTCTGCAG GTTGAACCTGG TTTTCTGTT TCTTGATTG	3660
AGGTACCTTC TACTGTGCCT TCATTTGGAT TTACTGGAAC TTCTTCTACA GTTTTTCTG	3720
AATTTCATT TTTAGAGTCA TTATGTTCTG GTTTATTGAT TTCTCCAATC GAGGTTGTG	3780
AATCACTAGG ATTACTGGAC ACTTCCCCAG TATTTTGCT AGATGTATCT GGTGATACTT	3840
TCTCTGAATT CGTTGTTGAT TCTTCTGCAG GTTGAACCTGG ATTTTCTGCT TCTTGAAATTG	3900
AGGTTCCCTTC TGTAGTACCT TCATTTGGAT TTACTGGTGT TTCTTCTGTT GGTTTTACTG	3960

299

GAACCTCTTC AGTTTTTCT GGACCTTGTG CTTGGTCTT CTCAACCGGA GTTCAGGTT	4020
TTACTTGCTC AATATTACCC TTATATTCTG GAAGCGGTGC TACCTGCTCT GGTCACCTT	4080
TATCACTTAC CACAGTATCT GGCAGCTCTG GTTGAACCTC AGTCTCACCT TTGTCGGTCA	4140
CAACTGCTTC GGGTAATGTA GGTTGAACCTT CTGGTTCGCC TTTGTCACTT ACTACAGCTT	4200
CGGGCAACTC AGGCTGAATT CGGGGTTCAA CAATAGCTCC AGACTGTACG TCCTTATGTT	4260
CTACACCAAGT CTCAGGTTGT TCCTTATAA CTTGAGTTTT TTTAGTACCT TTTTCGACTA	4320
TTCTTGGACT AGGCGCAGTC GTTGAAGTTG AAACAATTTC TCGCGAAACT TCTTCCTTGT	4380
TTACAGAGAA TATTCTGACG ATTCAACTT TCTTACCTAA TTTACCTTCT TGTTTTACTC	4440
TTACAGTTCC TTCAGCTAAA TCAGGATTTT CTTGAATTTC TTCTTGAAAA TCTATTTTG	4500
TCTCCATAGT TTCCCTCACGA TATAAGAGTT CAGGTTCTT CAATTGACCT GATAAAACTT	4560
CATCCTGTGG ATTTAATGTA TTTACCCAG TCTTTCTTT TGGAGAAATC TTCTCCTCTT	4620
TCTTCGTTTC TAGATTCTTA TGTCGGCTA ATTGTTCTTGA AGAATCTGAA GATTGTTCT	4680
CTTCTTTCT TGGATGTGATT AATTCACTGAG AGAAAGTTT TTCAACTACT TGAACTTCTG	4740
TCGGCTTAGT TGAAGAAACA GGTGTTGTT CCTGAATAGC TTGTACTGTT GATGGATGGT	4800
CTACAAAATT CGGTGTAACA TTATAATCCA CCTTTGTTG TTTGTAGGA GTGGCAACTG	4860
AACTCTTTG ATTACTTACT TCAGACTCAG AAGTCGTTT TCCCTTTTG ATATATCCAA	4920
TATAACTGTA ACCTGAAATC TCTTCTGAA GAGGTAATTT TTCTCCAGAG GTCAATTCT	4980
AGTCCGTATT GTAATTAGC AAAAGATGAT TTCTAAAGC ATGGACTGAA ACTAAGACAC	5040
CATTTCTTAT CCCTGCAACC AATACTAAAT GTAAATACCGT TTTATTCTTA ACCTTTTCT	5100
TGGAAACAGC AAAAATTAAA ATTCCCATAG CAGCTAAGCT AGCACCAGCA ACTAGGGCTT	5160
GCCTCTCATT CTTGTTCCA GTATTTGGCA ATTCCGCCAG TTGATTTGA GAATTAACT	5220
TATAAAACAAG ATAATAAGTT TCATCATCAT TCTCCACGTA TGTCGGAATA TCATAGACAA	5280
GCTGCTTCTT TTCTCTGAT GATAGCTCTG AATCTGCCAC ATATTTATAG TGAACTCCCG	5340
CAGTTTCTTGCAGA TACAAGTCCT ACTGATAATT TTCTAAATGA AAAACGCTCT TGTTTTAC	5400
TCGTTGCAGA TACAAGTCCT ACTGATAATT TTCTAAATGA AAAACGCTCT TGTTTTAC	5460
CAAAATACCTT TTCCATTATT CCTCCTTGAA ATAAAATTAA TATATGTTAC AAAGACCTTT	5520
ATTATATTAG TGTATTATCT ATTATCTATA GAAAAGGCAG TATACCTAA TTATACTCTT	5580
AATTTACAAA AAAGTCTTAA AATTGAGATG CGCTTTCATA CTTTGTGTTA TATTATTTGG	5640
AGGTACAATA ACACCTACCA TGAAATTTAC ACGGTAGGTG TTACTCATAT CACTAATCGT	5700

300						
TCTAAAAATG	GTTGAGGCA	GTTGAGGAGA	ATTCCTTCTA	TCCAGCTTCC	TTGTGCTGAT	5760
GAGCGATGGT	CTTCCTGCAG	GCTTTTTTTT	AGAAAATCTC	GGACTTGTC	TGGTGCAGATT	5820
TCAAATTCAA	AGGCTTTCAT	TTTATAGAAA	AAGTCGATGA	GATGATCTGA	CAGGTATTCA	5880
GTTGAAAAGG	GTACTTCACC	ACTTTTTCTA	TATTCTAATA	AGAGTCTAGA	AAATCGAGCT	5940
TTTTCTTCAG	GAAGCTCACG	AAAATAGGAA	TTGAGGATCC	AAGTCTGCTT	CTGTTTCTT	6000
TCAATTGGAT	CCTGACTGGC	AATTCTGG	TCTTTTCCA	GCTTTTTTG	GTATTGTTG	6060
GCCTTGATAG	CTCGTTCTGC	TCTATTTTA	CCAAAAGAA	TTTTTCCA	CTTGCCTTCT	6120
TCTTGAGTCA	GGGTCTCTGT	AAAGCCAAAG	TAATCTTGAT	AAGCACGCTC	TGCGGGTCCC	6180
ATGGCTAGAA	CCAGATTGTC	TGCATATTGC	TTGGCGATTT	TATCCCTCTT	CTTGCCTTCT	6240
TTCTCTGCCT	GGATAACGGAG	TTCTGTTCG	TAGTCATA	TCTCCTTGCC	TAGCTTGACA	6300
AGGTAGAGTT	GGTCATCCGA	TTTCCCAAGT	AAAAAGGGTT	TGATACACTT	TTCAAGGACT	6360
TCTTCCATCC	GAGCCTTTT	CTTGGTTCC	GCCTTGGTCC	AACTCCCTCC	CTGAAAGACT	6420
TCTAGGAAAA	GCTGGTAGTC	TCTCTCAGGC	GCAAATTGAT	TGCCACGATT	GGGTTTGAAA	6480
ACACCTTTT	CCCAGAGCCA	TTTAAAGAGT	CGCTCGTCAA	AGTTACTTTT	ATTGACCTTG	6540
ATTTTTTCCT	TTTCTGAGC	TTTCTGGTT	AGATTTCAA	CCTTTCTGAG	CAGTTTTCT	6600
TCCTCTCCA	ATTGCTGGTC	AAAGGGACAAT	CGATGAAAAT	GACGAACACA	GTCGCTACCA	6660
ATTGGAAAGA	GGCGTTGCC	TGTGACACCG	TTAAAGAGTT	CATAAGCGTA	TTTGATGGCA	6720
TTTCCACAGA	CACAATTGCT	ACGGCCGATA	CCGTTAAAAA	TAAAGGAAAC	TTCAATTCCAT	6780
TCCTTGGTAG	CTTGGTCCCA	AGTATCCGCT	TTCGAAGCCT	GTAAAACCTGC	ATCGTGCAGG	6840
GATTTTCTAA	CTGGAAGTGT	CATGAGGTCT	CCTTTCTAAAT	ACTCAATAAA	AATCAAAGAG	6900
CAAACCTAGAA	AGCTAGCCGC	AATCAGCTCA	AAACACTGTT	TTGAGGTTGT	AGATAGAACT	6960
GACGAAGTCA	GCTCAAACAA	CTGTTTGAG	GTGTTGGATA	GAAC TGACGA	AGTCAGTAAC	7020
CATATATACA	GCAAGGCAGA	GCTGACGTGG	TTTGAAGAGA	TTTCAAAGA	GTATAAGTTA	7080
TACTTTTACA	ACTGAAACCT	CGTCTTCTACC	GAGTAAAATC	AAGTATTTT	CAATATTTTC	7140
AATCGAATAG	GCTCGTGATA	AAGCCTTTC	GTATAGAGCT	AACTGACCAC	GATAGCGGTC	7200
TACGAGTTGA	CTTGGTTCAT	CATAGCGGTC	TGTCTTGTAG	TCGAACAGAA	CAATTTTGT	7260
TTCGTAAAGC	AGATAGCCAT	CAAGGATACC	ACGGACAACA	AACTCTTCT	GACTCTTTG	7320
GTCTCGTTG	AGCATGGAGA	AAGGTTGCTC	GCGATAAAAGA	TGGTCGGTAT	TAGCAAGAAT	7380
TTCCTGACCG	AGTACTGTGT	CAAAGAAAGC	AAAATTTA	TCAAGATTGA	TCTTGTCTCT	7440
GACAGCTTGG	CTAGTTGAA	CTTGGTTGAG	TGTTTCTGTT	AGGCTAGCAA	GGGTTAGTTG	7500

301

CTGGCTGAGG TCAATTCTCT GCATGAGTTC GTGAGTAGCA CTACCAATCT CAGCTCCAGT	7560
TACCTTTCT TTGGTGAAA AATCTGGCAA ATCGAAGCTG ATTTCTTGC CTACTGACTG	7620
ACCTTGACCA GCAATCTCGA CACCTTCCAT ATCCATAACT GGTCGTAGA ATTTCTTGAT	7680
TTGACTTGGG GTTTGAACAC TAGGAAGTTC AATAGCTGCG CGGTGAAGAG TATTATAAAC	7740
TTCCACCTCC TTCAGCATTT CCAGAGCTTC TTGATGGTA TCTGACTGAC GATTGTCTGC	7800
TTGGGAGCTA TCTTGGAGAG GACTCTGGT TTCCAACTCT CCGATAGCTT CTCTGGTCAA	7860
CTGATCTTCG CCAATAAAAC GATAACTAAA GTTGAGCTTG TCCTTAGTAA ACACTTACT	7920
GATAGCCAA AGCCAATCTT GGAAATTCCG TGCTTGCACT CTAGTATTGC TATTTAGTTT	7980
CCCATTGGT GCTGCTGGGT ATTCCTTGGA TTCCAGCTT TCACGAGAAC CCTTGCCGAC	8040
AAGATAGAGC TTTTCTCA CGCCGCGTCAT AGCAACATAC AGCAACACGCA TCTGCTCAGA	8100
ATAGCTTGCT AGCTGTAATT CCTCTTCGTT CTGCCTATAG GTCAGACTAG GAATGGAGAG	8160
TTTGATGGTT TTAGGATAGT GGTCTTCTAC TGCCCTGTC TCCATCTTGG CAATATATTT	8220
GACACCAAGA CCATTCTGAC GACTGAGAAT GACTTCTGAC ATAGAGTCTT GCTTGTGAA	8280
ATCTTGATCC ATATTGAGGA TAAAGACGTA AGGAAACTCC AGCCCTTTAC TCTTGTGGAT	8340
GGTCATGAGC TCTACTGCAT CTTTGGCGG TGCGACGGCC ACGCTTGCCA AATCGTGTG	8400
GGCTTCTAAG ACTTGGTCAA TCATACGAAT AAAACGCGAC AAACCTTGA AATTGCTCTT	8460
TTCAAATTGA TCAGCACGCA GTGCTAGGGC ATAGAGATTG GCCTGCCTAG CAGGACCATT	8520
CGGCAAAGCC CCAACATAGT CATAATAAA ACGGTCGTTG TAAATCTTCC AAATCAAGTC	8580
ATAGAGAGAG TGGTTTTGG CATACAAGCG CCAAGAAGCT AGGATATCCA TGAATTGCTT	8640
TAGTTTTCA GCTAGAGCTG TGTGAATCAA GCCTTTTGTA CTACTTGCCA TTTTTGTGC	8700
ATTGACCAAGT TTCTCATAGA GATTTCTGTC TTTATCC TCTGCTTCT GAAGGGACAA	8760
ACGTGCTAGC TCATCCTCAT CAAAACAAA CATTGGAGAC TTCATAAGGG CAACCAAGGC	8820
GTAGTCTGC AGGGGATTGT GAATGACACG AAGAGTGTCT AGCATGACTT GCACTTCTAG	8880
GGATTGGAGA TAATTGTTT GCTCTCCGTG ACTTTGACA GGAATTCCGT ACTCAGACAG	8940
GGCGAGGAGA ATCTGGTCAT TACGACTGCG GCTGGAGGTC AGAAGGGCAA TTTCCTTAAA	9000
GGCAACACCT TTTTCTTGAT GAAGTTTCAG AATCTCCTTG ATAACATAAGC GCATTTCGCC	9060
TGTTAGTTTC GTTCTGTTT GACTCTCTTC TTCCACCT GTATCGTCT TGCGTAGAG	9120
GAGAAATGCT GCCTGTTGT CTGGATTGGG AGTCAGTTG GTATTGGCAA AAACAAGCTG	9180
GTGCTTGTAA TCATAGTTGA TTTCGCGAC CTCTGGTCC ATGAGACGTT CAAAGACATC	9240

302

ATTGGTTGCT GACAGCACTT CTGAACTACT ACGGAAATTTC	9300
GCCTTCTGG GGATTTGCG CATAGCGTTG GAATTCTCA TTGAAAATCT	9360
CTGACGGAAA CGATAGATGG ATTGCTTGAT ATCTCCCACC ATAAAGCGAT TGTGGCCATT	9420
AGACAACAAT TCCAGCATCC GTTCTTGAAT ATGGTTGGTA TCCTGATACT CATCGACCAT	9480
GACTTCATGG AAGCGCTCCT GATAAGACTC ACGAACTTGT GGGAAATTCTC	9540
AATGGTGTAA TGGCTGATAT CAGCGAATTC GAAGGCATTT TCCTGTCGTT TTCTCTGACG	9600
ATAAGCCTCT ACAAAATCGC TCATGAAAGA TTGGAAGGTT TTAGCTAGTT TCCAAGTGT	9660
TCCATGATAA CGTTCTTGAT AGTCGAGAAT CGCTATCTGG TCTGATAATT GTCTAGTTT	9720
AGCAAACCTGG GTCTTCTCT CTTCGTTGTA GGCATCAGCC AGGGGCTTCA AATCAGCCTA	9780
CGGCTGGCAT TAGTCAGAGC TCGACCGTTT TTCTCCTTAG AGATGGCGAC AACACGGCA	9840
AGCACTGCCT GATAAGCCTG ACTATCGGAC TCCTGATTTA GGGAGCCAAT TTCATCCAGA	9900
ATTAACGTAA CATTCTAA ATAGGCAGCC TTGCAAACT CCTTGGCATC GTTATCCAGA	9960
TGGTAACGGA AAAAGCTTTC CAAATCCAA AGGGCTTGTT TGATTTGCTC GGTCAGTTT	10020
TCTTTTCAC TGGTAAATC AGCTTCTCA AATCCTTGA GGAAAGATTTC ACTCAGCCAC	10080
TTTGAGGAT TACTGGTGA TTGGAGGAAG TCATAGATT TATAGACCTG CTGGCGCAGA	10140
CCCCGTTCGT CCTTGCACG CCCAGCAAAG TTTTCAGCA AATGACTAAA GGTCTCTTC	10200
TGTTTACCTT GGTAATGCGC TTCAAAGACC TCATGAAAGA CTTCGTTTC GAGAATAAGT	10260
TGCTCGCTT GGTTTGTA AATACGGAAA TTAGGTGCAA TATCAAGCAG ATAACCATGT	10320
TTGCCAAGGA ATTTTGTTGT GAAAGAATCC ATGGTTCCAA TGGCAGCGTT GGGTAGGTCT	10380
GCCAACGGC GACCCAAGTG TTGTTTGAGG TCGACATCAT CTGTTCTTG GATTTCTTG	10440
CTGATTTTT TCTCTAAACG TTCTTAAAGT TCAGTTGCAG CCTTGACGGT AAAGGTTGAG	10500
ATAAAGAGTT GAGAAATTTC GACACCACGC GCCAATTGGT CCAGAATGCC CTCTGCCATG	10560
ACAAAGGTCT TTCCAGAACCC AGCCGATGCT GAGACCAGGA TATTCTGGGC AGAAGTGTAG	10620
ATAGCTTCGA TTGCTCGGC AGTTTCTTC TGTTCTTGC TCGAATTGCT TTCTGCTTCT	10680
TGCAGTTTT GAATCTCCTC CTCACTTAA AAGGAAATAA GCTTCATCGA TTCAACTCCT	10740
CTCTTATTT TTCAAGCCAA GCTTGCTTGA GTTTTCTCC GACCAGACGC TTGCCATCAG	10800
CTAGGTCCAA CTTTTCTAGG AAACGGGCTT GGCCCAGATG GTAAATTGGCT TCAAAGCTG	10860
TAATAGCCTG ATGTTGCTGG ACGTATGGGG CAATGCTTCT GCCATTTCA GTATAAGGAT	10920
TGATGGCGAA CCGGCCTGCT AAAATCTTCT CAGCAGCTTT CTTGTAAAGA TAGGCATTGT	10980
AGTCCAGTAG GAGCTGAAAT TCCTCATCTG TCAGTTGATT AGCCTTGTGTT TTGTTATAAA	11040

ATTCGCCCTAA ATAAGTGCTT TCTTTTCCA AGAAGAGCCC TTGGTATTC ATAGATTCG	11100
TGGCTCTAC CACTGCTCCT GCCAGACTTT TTACCGCCAT CAGAGATTGG ACAGGTTCA	11160
CCATTTCCTAA GTACATGGCG CCGAAAAACT TCTGCTCCCC TTCTCTTTTT AGGGCAGCAA	11220
GATAGGTTGG TAACTGAGAA TTGAGCCAT TAAAGAAATG AGGAAACTGG AACTGAGTCA	11280
GACTGGATTT GTAGTCTACT ACTCCTATCG CTCCATTAGC TTTCAAAACGG TCAATCCGGT	11340
CCACCTTGCC TCGTACAAAG ACACTGCGTC CATTGTCTAA TTGAATAAAG GCTTGGTCTT	11400
TTCCACCAAA ATTTGCTTCT TCTTTGATGG TTTCGATGGC TGGATTGTGT CGGAGAATAT	11460
GTCCAGTTGT CCGTCAACA TCAAGCAAA CTTCCCTGGT AAACCTGGCT TCCAAACTTT	11520
CTTGATAAAT AGCTCAAAT TCGCGTTCTT GACTGGTTTC TTGAATAGCT TGTTCAGAC	11580
GTTGGTCAA GGAATCTTCA TTAGGCAACT GTAAGGCGCG TTCAAGATA CGATCCAAGA	11640
AATTCCCGTG ACTACGGGCA TCAGGATGCA AACGTAATTG CTCCTGCAAG CCTAAACGT	11700
AGCGTAGGAA ATAAGTGTAT TCATTGCGAT AAAACTCTGT CAAACCCGAC GTAGACAGGT	11760
AAAACCTCCTG TTTGGCAGGA TAGAGAGCTT GCAAGGTGTC CTTGGCTAAG GTCTTGCTGC	11820
TTGGACTGGT TGGGATAGCT GGATTTCCA GACCTTGCTG ATCTAGTTTT TTACCTATGA	11880
CACGCGACAG AACCTTGACA AAAAGTCAAAT CTTGCTCAGT ATCGCTCATC TCACCCCTGCT	11940
GGTGATAGGC AACCAAGACTA GACAAAAGAC TGTGATAGGA CCCCATATCC TCCTTAGACA	12000
GTCCTTCTC ATTCACTCTC TTCTCTCTCC GCCTAAATCC AAAATGGATC AACCTCTTGAA	12060
GATAGGCAGA TTCCTTACTT TCACCTTCGT TAAAAAGGCT TGGAGCCGAC AAGAACAACT	12120
GCTTACGAGC AGAATTGACC AAGGAAAGCA TAGTGTAGCG ATTTTTCTTG AGATTTTAC	12180
TGCTGGCAAT CAGTAATTGA ACGCCTTCTT CGGTCGCTTG GTTTAGGTTT TGCCCTTCTT	12240
CATCTGTCAG AAGACTGGTG TTTTGAGAAA TTTTGTTGAA ATTGTCCCTGA GTTAGTCCAA	12300
TAGCATAGAC AAAGTCAGCA GTCAATGGTG CAATCAAATC GTAACTCTGC ACCAGAACAG	12360
TGTCCACTGT TGCTGGAATG GTACGGTATT GGGACAAACT CATTCCAGAA TGGAGCAAGG	12420
CTAGGAAGTC TTCCAGACTA ACCTGTGAAC CAGCAAAAC AGTCGCAAAT TGTTCTAAA	12480
CATGGCAGAA AGCCTTCCAA ACTTCGGCTT GTCTTTCTTG TTCTACAGCT TCCAAAGTGG	12540
TTGTCAAATC TTGTAACTGC TTGGTCACAG CTCCTTCTTT TAGAAAGACA CTCCATTTC	12600
GTAGGAGTTT TTCAAGCCTTT TGTGTTCGGC TGGCAAAGAG GGTTCAAGA GGTGCTAAA	12660
TTCTCAGGCG GAGGACATTC AAACGCTCAA GATTAATTT TCCATGGTGG GATTTGGTGA	12720
AGGTTTGCTG AAAGGCTGGC AAGCCATTGA TACCAAGATA GCGGATATAT TGCTAAAAAG	12780

304		
CATCAATATC	AGACTGACTG	AGGTCAGTAT
ACAAATCACT	TCTAAAGAAGA	TTAATCAAAT
CCTCCTGACG	AAAACGGTAA	CGTTTTAAAG
CTAAAATAGA	CTCGACAAAC	TGAGTCAGG
GATGATGAGC	CATGGCTTCG	CTTCTACCAA
GATAAAAAGG	AATCTGATAC	TGGTCAAAA
TGGTTTGAG	AGATAACTGG	TAAGAAGCTA
CATCCCCAA	GAGAAATACGA	AAATGCTTGT
AGCTCAGGTC	TGAGTTCTCA	TGTAATTCT
GACGAATACT	ACGGGCTACT	AGCTCCAAC
CCTCCTTTG	CGTCACAAAC	GACCAGATTT
GTAATTTTC	ACGGTCTTC	TCATCGACAT
CCAAAGCGAG	TTCTGAAAAG	TCATAAGAAG
ACTCCAACAA	ACGAGAGGCC	TTGTCAAAAC
TATCCATCTT	CTCATGAGTT	TGAGAACAGT
CCTGAGCAGG	CGTTGGTAT	TTAGAAGCCA
GATGATGGAG	AAATTTACG	CTGGCTTGGT
AGAGATTGCC	CTCGCTAAA	GGACTGGTAT
AGGCTTCTT	ACTAGCATAA	GCCCCGATAA
CAATCTCAAC	ACCTTTGCCG	TGAAGTAAGT
CCACAAACCG	CTCTCCCTCA	GCAGAAAAAC
GAGTAAAGCC	GTCAATGACC	AAGGCCATT
GATTAAAATC	ACTACTTACC	TTGTCATTCT
CAATAGCCTC	AAATCAAATGG	GACAACGTGAC
TTTCCTGGGC	TAACTGACCT	TGATTAAGAT
AGGCTGTTAC	TTTCTCAAA	ATCAAGAGTA
AATCCGCCCT	CTTATCCTCA	TCTGTTAAAT
TCTCCAAGTC	CAAAAAACTC	ATCTGAGATT
TGGTCATCTC	ATGGTAAAGC	TCAATTAAC
GCTGGATCAA	TTGAGGATCC	TGCTTAATAG
CGCCATAAAAC	ACGCAAGTCC	TTGGGATCGA
GTTCGGCAAG	GCATTTGTAA	AAGGCCAAC
CAAGACCGAT	ATCATCAAGA	GTAGTTTAG
CTGGTAAATC	ATTCAAGACC	AGATAGCGAG
CCATTGAGC	AAAGCGCGTG	ACGTAATCG
AAAAAGAACG	CTGCTGGAC	AAGTATTCCA
GCACGGCGCG	TTCCCTTTCA	AAAGAAAGAG
AGTTGGGGGC	AATGTAGAAG	ACCCGCTTGC
CAGCTGCAAC	TAGCTCTTCT	GCCTCTCTTG
TTAGAATTTC	TGTCAAAGAA	GTCCGAATAT
CAGTATAAAG	TAATTCATC	TCAGCCTCGT
TGGAATTTTT	CATCACCCCTA	TATTATACCA
TGATTAGCCT	CGTAAATCTG	TTAAAATATT
TAGGCCATCC	TTTCTTTCT	TCATCATCTG
CTAAATCTTA	AATACTTAGC	TTTACTTGTA
TTAGATAGAA	TAAGTCTGGC	TACTGAAAAT
CACATAATAA	AAAAGCCTCG	GTAAACAAGGC
TTTGAGTTT	ATGATTGT	CTTAGGTACG
GAATACACTT	CAATGTGTTG	TCCCAGTATC
TTAATGTCGA	CTGGTAGATT	GTCTGATT
TCGCCATCAA	CATCGGACTC	TAATTCGATA
TCAGAAGAACG	TTTTAATATT	ACGTGCCTT
ATATATTCAA	TATTCTTGAT	AGAATGATTG
AACTATAGTA	AATTGAAACT	ATAATAGTAC
ACCGTGGATG	CTAAAATATT	TCTAGAAATT
AATTGATT	CCCTAATCAA	GCTATTGTA
TCTTATTTC	ATCTACTATA	ATAAAATGAA
CCAAAAATAG	TACACAATGT	GGTATAATCT
TCTTATGGCA	TATTCAATAG	ATTTTCGTAA
AAAAGTTCTC	TCTTATTGTG	AGCGAACAGG
		14580

305

TA GTATAACA GAAGCATCAC AC GTTTCCA AATCTCACGT AATACCATT ATGGCTGGTT	14640
AAAGCTAAA GAGAAAACAG GAGAGCTAA CCACCAAGTA AAAGGAACAA ACCAAGAAA	14700
AGTGATAGA GATAGACTTA AAAACTATCT TACTGACAAT CCAGATGCTT ATTTGACTGA	14760
AATAGCTTCT GACTTGGCT GTCATCCAAC TACCATCCAC TATGCGCTCA AAGCTATGGG	14820
CTACACTCGA AAAAAAGAAC CACACCTACT ATGAACAAGA CCCAGAAAAA GTAGCCTTAT	14880
TTCTTAAGAA TTAAATAGT TTAAAGCACC TAGCACCTGT TTAGATTGAC GAAACAGGAT	14940
TCGATACTTA TTTTATCGA GAATATGGTC GCTCATTAAA AGGTCAGTTA ATAAGAGGCA	15000
AAGTATCTGG AAGAAGATAT CAGAGGATT CTTTGGTTGC AGGTCTAACAA AATGGTGAAT	15060
TAATCGCTCC AATGACTTAC GAAGAGACGA TGACGAGCGA CTTTTTGAA GCTTGGTTTC	15120
AGAAGTTTCT CTTACCAACA TTAACCACAC CATCGGTTAT TATAGTAAAA TGAAATAAGA	15180
ATAGGGGGGG GGGGGGAGGG GGGGGGAGGG AGA	15213

(2) INFORMATION FOR SEQ ID NO: 27:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6004 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 27:

TTATTACCTG AACATTAAA TTTAATTGGA CATCCGTTA TCAATTAT TATCATCA	60
AGATTTTAT TATCTGATTC AGGAATTAA TCTGATATAA CAACACCATT TTCAAGATAG	120
TTCATTAAAT TATTTGATTC ACTAACATTA GTGTTTGAT CTCCATCAAG CAAAAATAA	180
TGGTTATCGG AACTAAATA CGATGAGTTT AAAATTAT TACAAATTAT TTGATTTGCT	240
CCACCAGGAA TATATCTCAC TACTAAATTG TGTAAAGAT TCTCACTACC TGAATGAGTG	300
ATAACAAACT CTAGAATATA TTTAGCTAGT CTATCTCAA CATAAATCAT CTTCTAGAA	360
TGATACACAT CACCTAATTG AAAAAATGCA CCCTGATAAT CAATTTTC AATAACATCT	420
ACCTTTCTC CGTTTTTCAC TAAAGTTTC ACGGCTTCTC TAGGAAATC TTTTATAAGT	480
TGTGTAGAAT GTGTAGTGAT AATAATTG A TGTGTTTAT TTAAACACTC TTGAAGTAAA	540
AACTCTTAA ATTTATAGAT TGCACTCGGA TGAAGTGAGA TTTCAGGTTTC ATCTATTAA	600
ATTAATGAAT TTGATTGCGC ATTTACTATA TCATTTACTA ACAAAATAAT TCTAGCCTCA	660
CCTGTTCTG CAAAAGCCTC GGAATATTCT TTTCCAGATT TTTCATCCA AATAGTTTG	720

306	
GAAGCTTTA TATCATCACC TTTGAAATAC AACTTATGTG TTAATTTG AATGTCTGTA	780
TAAGATTCAT CCATTATTC ACTAATAATT TCACAAACTT TATCATCAAC TTTAACATTA	840
TCTATAACCA TTTCTTTT ATAACCGGTA TAGCTACTTG TATTATTCTT TAAATATCA	900
GCAACTGGCT TAGATCGTAA TCTTATAAAA TCTTGTTCAC TACGTTGAGT AGAAATTTT	960
TTAAATATT AGTGATAGAA AAATAATCA AAAGCAGAAA CATATTCTT ACAATCACAA	1020
AAGACAAACAT TTTTTCAAT GCCATCCCAT CTGCTGTG AAGAACTTCC AATATATTAA	1080
TTTTGGGTA ATCTTCCAT CTCATATTGT TTTTGAGGAG CATATGGTTC CCAATAATCT	1140
AATCCTTTT TTGTTCCAGA ACGGCCTTA AGAACTTCTA CATTCTAGA AGCTTAAATG	1200
TTATAATATG AATAGATTAAC ATATTGTTTC CCATCCACTT CATCTATTG ATCAACATTT	1260
GTACTAAACC AATATTCAGA CACACTTTA TTGGCTGGAG AACCATATAA AGCTTGTAAA	1320
ATTGAAGTTT TATTACTCC ATATCTATTA CAGACACCTC AGGATTATTT AACTTATAAG	1380
TTTTAACAGC TACGGAATCA ATTTCAACAG CAACTTGAAC ATCTATGCCT GATTTTTAA	1440
GGCCACTTGT AGTGCCACCT GCACCGTTAA ATAAATCAAT AGCAACAATT TTCCCCATAG	1500
TATTCTCCTA AAGTTCTCC TTTTATTAT AACATTATCA AATGAAAC CCAACCCGAT	1560
AGGGTTAGGT TTTAACATC ATTTCACCAA CTTCTTCATC TCATCAATAC GTGCGACGGT	1620
CGCGTCATAT TTAGCTTGAGT AGTCAGCTTG TTGTCGAT TCTTTGGA CGACTTCTGG	1680
TTTGGCGTTG GCTACGAAGC GTTCGTTAGA GAGTTCTTA CCAACCAGT CCAGTTCTTT	1740
TTGCCATTAA GCAAGTTCCCT TGTCGAGACG GGCCAGTTCT TCTTCAACAT TGAGGAGATC	1800
GGCCAGTGGC AGGTAGATTT CTGCTCCTGT GATGACACTT GACATAGCCA GTTCAGGTGC	1860
AGGGATGGTT GATGCGATTT CCAAGTGTTC TGGATTGTA AAGCGTTGAA TATAGTTGAC	1920
ATTGCTGTTA AAGAAGGCTT CCAAGTCGCT ATCGTTGTC TTAACAAGGA TGGTGATAGG	1980
CTTGCTTGGT GCTACATTAA CTTCCGCACG CGCATTCCGA ACAGCACGAA TCAAGTCTTT	2040
GAGACTTCC ACACCAAGTGT GAGCCGCAAG GTCTCAAAG GCTAGATTAA CAGTTGGGTA	2100
TGCAGCTGTC ACGATAGAAC CTTCTGAGAT TTGTCCTAG ATTCCTCTG TCACGAATGG	2160
CATGATTGGG TGAAGGAGAC GAAGGATCTT GTCCAGCGTA TAGAGGAGAA CAGATCGAGT	2220
AATGACCTTA TCGCTTCAT TGTCGCTGTA TAGAACTTCC TTGGTCAACT CAACATACCA	2280
GTTGCCAAAT TCTTCCCAGA TGAAGTTGTA AAGGATATGA CCAGCCACAC CAAACTCGAA	2340
CTTATCAAAG TTTTCAGTAA CTTTGCAAT GGTTCGTTG AGATTGTTGAA GAATCCAGCG	2400
GTCCCGTCACA TTACCAAGCCT CACCTGTTGC AACTTTGAG ACATTGTCAT GCGCCACATC	2460
CAGCGTCAAA CCTTCATTGT TCATGAGGAT ATAGCGAGAA ATGTTCCAAA TTTTGTAAAT	2520

AAAGTTCCAT	GAAGCATCCA	TTTTCTCGTA	AGAGAAACGA	ACGTCTTGAC	CTGGTGCAGA	2580
ACCGTTTGAA	AGGAACCAAC	GAAGGGCATC	ACGACCGTAT	TTCTCGATGA	CATCCATTGG	2640
GTCAATCCCG	TTACCGAGAG	ATTTAGACAT	CTTGCCTCCT	TGCTCGTCAC	GGATGAGACC	2700
GTGGATAAGC	ACGTTTGGA	ATGGCTGACG	ACCAAGTAAAT	TCCAAGGACT	GGAAGATCAT	2760
ACGAGACACC	CAGAAGAAGA	TGATGTCGTA	ACCTGTTACC	AAGGTTGAAG	TTGGGAAATA	2820
ACGTTAAAG	TCTTCTGAGT	CGACTTCAGG	CCAGCCCCATG	GTTGAAAATG	GCCAGAGGCC	2880
AGAACTGAAC	CAAGTATCCA	AGACGTCCTTC	GTCCTGAGTC	CATCCGTCAC	CTTCTGGAGC	2940
TTCTTCGCGG	ACATACATTT	CACCATCAGC	ATTGTACCAAG	GCAGGGATTG	GGTGACCCCC	3000
CCAAAGCTGA	CGAGAGATAA	CCCAGTCGTG	GACATTTTCC	ATCCATTGAA	GGAAGGTATC	3060
GTTGAAACGA	GGTGGGTAGA	ATTCGACCTT	GTCCTCTGTG	TCTTGGTTAG	CAATGGCGTT	3120
CTTAGCCAAT	TGGTCCATCT	TGACGAACCA	TTGAGTAGAC	AAGCGTGGCT	CAACTACGAC	3180
ACCTGTACGT	TCTGAGTGAC	CAACACTGTG	GACACGTTT	TCGATTTGA	CAAGGGCACC	3240
GATTTCTTCC	AACTTAGCAA	CGACTGCCTT	ACGAGCTTCA	AAACGATCCA	TGCCTGAAAA	3300
TTCAAAGGCA	AGCTCATTCA	TAGTTCCGTC	GTCGTTCATG	ACGTTGACTT	GTGGCAAGTT	3360
ATGACGTTGG	CCAACCAAGA	AGTCATTTGG	ATCGTGGGCA	GGTGTGATTT	TCACGACACC	3420
AGTACCAAGC	TCAGGATCTG	CCTGCTCATC	TCCAACGATT	GGGATGAGTT	TATTAGCGAT	3480
TGGAAACGTC	ACCTTTTAC	CAATCAACTC	CTTGTAGCGC	GGGTCTCTG	GAATTAAACCC	3540
AACCGCAACG	TCCCCAAACA	TAGTCTCAGG	ACGAGTTGTA	GCAACTTCAA	GGGCAGCGTGA	3600
ACCATCTTCC	AGCATGTAA	TCATGTGGTA	GAAGGCACCT	TCTACATCCT	TGTGAATCAC	3660
CTCAATATCA	GAAAGGGCTG	TGCGAGCTGC	TGGGTCCCAG	TTGATGATAA	ACTCACCACG	3720
ATAGATCCAG	CCTTCTCTGT	AAAGGTTCAC	AAAGACCTTA	CGAACAGCTT	TTGACAAACC	3780
TTCATCAAGA	GTGAAACGCT	CACGAGAATA	GTCTACAGAA	AGCCCCATCT	TGCCCCATTG	3840
TTCCCTGTATG	GTAGTGGCAT	ATTCGTCCTT	CCATTCCCAG	ACCTTCGTCA	AGAAAGACTC	3900
ACGACCTAGG	TCATAACCGG	TAATACCCTC	ACACGTAAG	CGCTCCTCAA	CCTTAGCCTG	3960
AGTCGCAATA	CCAGCGTGGT	CCATACCTGG	AAAGCCAAAGG	GTATCAAAGC	CTTGCATGCC	4020
TTTTGACGG	ATGATGATAT	CCTGCAAAGT	CGTATCCAA	GCGTGACCAA	GGTGAAGTTT	4080
CCCAGTTACG	TTTGGTGGT	GAATCACCGAT	TGAATAAGG	TTAGCCTTTT	GATGCCCTGA	4140
AGGCTTGAAA	ACATCCGCAT	CAAGCCATT	TTGGTAACGA	CCAGCCTCAA	CCTCGGCTGG	4200
ATTGTATT	GGTGAAGTT	CTTAGACAT	GTGTGTGTCC	TTTCTCTATT	TTGTTTATT	4260

308

TATTTTGAAAT	TTGCTTAGCA	GCTTCTTCTG	CAGACAAATT	CGTATTATTT	ATTTTAAAGT	4320
AGTGGTGCAA	CTCATTCCGGT	TGATGTTGGG	AATTTAATTG	AAGTGTTC	CGGGTCTCTA	4380
AAATTTCTCT	TTCAGATACC	TCAATATGTC	GTGTTAAGGG	TTTGTGCTTT	AATCGATTCT	4440
CCGTTGATT	TCGACGTATG	CACTCTCAA	GACTGTTC	CAATTCAACA	AACAGAATCT	4500
CTTGATGAAA	GTTATCCAAT	AAATCCTGAA	TTTGCTTTAA	ATACATCAGC	TGGTACTGAT	4560
TTGAAAAAATC	AATTACGTCT	GTGAAAATTA	CTGATCGCTG	ATTTCTTGCA	CTTGCTCCAA	4620
GGAAAGAAAA	GGTAATTCCA	CGAACAAATT	CCCACATCTC	CTCGGTATAA	TCCTGATAGA	4680
TCTCTAGTGC	AAAATCAATG	GCTTGATGGT	TATAAAATAG	GGTAGCATCC	GTCAGTCGAG	4740
ATAATTCTTG	ACCAATGGTC	ATTTTTCTG	ATGCTGGAGC	ACCAATGATG	AAAAGATGCA	4800
TCAAATCACC	TCCCACTCAC	TCCTCAGCAA	GCCATATCTC	AAATCATCAC	AGCAGTTGCC	4860
TTGAGCATCT	TTGGGGTCTC	TTATGCGAGC	TTCGAGGGTA	AAGCCAAGCT	TTTCCGAGAC	4920
TCGTTGACTT	TGAAGGTTAT	ATCCAAAGCA	AGTTAGTTCA	ATCTTGTGAA	GACCAAGTT	4980
TTTAAAAGCT	AGATCAATCA	AGGAACACGC	TGCTTCTGGA	ACATAACCTC	GACCCCAATA	5040
GTCTGGGTGC	AAGGTATAGC	CAAGCTCTAG	CACATCATCC	CGATGAAGAT	GGTTGAAGTC	5100
AACAGAACCA	ATGACTTTAT	CGGTTCCCTT	GACGACAATC	CCATAGCCAG	CTGGGAGATT	5160
TTCCTTTGA	GTACGCTCCG	GAAGAATGTG	CTCCAGATAA	AAATCTCAT	CTTCCAAGAT	5220
CTTGACTGGA	GGAAAACCTG	CTGGATAGGC	GACCTCTGGC	AAACTAGCGT	AGGTATGGAT	5280
ATCCTCAGCA	TCCACCACTG	TGGGACTCG	TAAAACGAGA	CGTTCTGT	TT CGATTTATC	5340
TGGCAGCTCA	GTTCTTGCCA	TCCTTCTTCC	TCGCTTTTTT	GATGAAACTG	CCCTTCATAT	5400
CTACACGCTT	GTCCAGATAG	CGATAAACGC	GCTGATATCC	ATCTCCCATG	AAATAGGTTG	5460
GGGCAAACAG	TTGATTTTA	AAATGTCCCT	TTTCATCCAG	GAGTTCTGGG	GCAACAAGTC	5520
GCTCAAGAAT	CTTGGCAAAG	ATGTGGCAA	TACCGTCTTC	CTCAACAATC	CTATCTACCC	5580
GACAATCTAA	AAACAGTGGA	CAGGCGTCTA	AAATAGGAGT	CTGAGTTCGT	TCAGAAATT	5640
CATAATGCAC	TCCCAAACGT	TCCAATTCT	CCTGATGACT	GATAAAACCA	GCCTGCTCCA	5700
TCGCAAGCAT	AGAAGTTCA	TCAGAAATAT	TCACAGTAAA	TTTTTGATAC	TGTTTGATCT	5760
GCTCTGCGGC	ATTCTCTCTC	GCAACGACTC	CAATCACAAAC	CCAATCTCCT	AGACTATAAG	5820
AGGAACTACA	GGTCGTGATG	TTATGCCAA	AATTCTAATC	TTGATATCCT	AAAATAAAAA	5880
CAGGAAAACC	ATAATATAGT	TTACTTGTGT	AAAAAGATTG	CTTCATAACA	ACCCCTTTG	5940
ACTAAGACGT	AAAAGAAAAG	CCCTGCCATC	TACATGACAG	GGACGAATGT	GTTTATCCGC	6000
GGGG						6004

(2) INFORMATION FOR SEQ ID NO: 28:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 5857 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 28:

TGTAGAATT	ACGACAATGC	TTCGTTGATT	TCTGGGTTGA	TTTCGTCGCG	TTCTGGCAAG	60
CGAGTCATG	AACCAAAAAT	AGTACACAAT	GTGGTATAAT	CCTTTTATGG	CATATTCAAT	120
AGATTTTCGT	AAAAAAGTTC	TCTCTTATTG	TGAGCGAAC	GGTACTATAA	CAGAAGCATC	180
ACACGTTTC	CAAATCTCAC	GTAATACCAT	TTATGGCTGG	TTAAAGCTAA	AAGAGAAAAC	240
AGGAGAGCTA	AACCACCAAG	AAAAGGAAC	AAAACCAAGA	AAAGTTGATA	GAGATAGACT	300
TAAAAACTAT	CTTACTGACA	ATCCAGATGC	TTATTTGACT	GAAATAGCTT	CTGACTTTGG	360
CTGTCATCCA	ACTACCATCC	ACTATGCGCT	CAAAGCTATG	GGCTACACTC	AAAAAAAGAA	420
CCACACCTAC	TATGAACAAG	ACCCAGAAA	AGTAGCCTTA	TTTCTTAAGA	ATTTTAATAG	480
TTTAAAGCAC	CTAACACCTG	TTTAGATTGA	CGAAACAGGA	TTCGATACTT	ATTTTTATCG	540
AGAATATGGT	CGCTCATTAA	AAAGTCAGTT	ATAAAGAGGC	AAAGTATCTG	GAAGAAGATA	600
TCAGAGGATT	TCTTGGTTG	CAGGTCTAAC	AAAAGGTGAG	TTAATCGCTC	CAATGACTTA	660
CGAAGAGACG	ATGACGAGCG	ACTTTTTG	AGCTTGGTTT	CAGAAGTTTC	TCTTACCAAC	720
ATTAACCACA	CCATCGGTTA	TTATTATGGA	TAATGCAAGA	TTCCATAGAA	TGGGGAGCT	780
AGAACTCTTG	TGTGAAGAGT	TTGGGTATAA	ACTTTTACCT	CTTCCTCCCT	ACTCACCTGA	840
GTACAATCCT	ATTGAGAAA	CATGGGCTCA	TATAAAAAG	CACCTCAAA	AGGTATTACC	900
AAGTTGCAAT	ACCTTTTATG	AGGCTTTTT	GTCTTGTCT	TGTTTCAATT	GACTATATAA	960
ATTGCTCTAAC	CGAAACAACC	GATAAGAATT	GGCACAAAAG	CGACCGTATT	TTTGTTACCA	1020
ATACAGGAAA	AACAGTTCAT	AGTTCTATCT	TGAGCAAGTC	TCTCCAGCGA	GCAAACGAAC	1080
GCCTTAAAAA	ACCAATTCCC	AAACATCTGT	CCCCTCACAT	CTTCAGACAC	ACCACTATTA	1140
GCATCTTATC	AGAAAATAAA	ATTCTTTAA	AAACAATCAC	GGACAGGGTT	GGTCATCCCG	1200
ACTCTGAAGT	CACTACTTCC	ATCTACACCC	ACGTCACAAA	GAACATGAAA	GATGAAGCAA	1260
TCAATGTACT	GGATAAAGTT	ATGAAAAAGA	TTTTTTAAAA	AGTTTTGTCC	CTTTTTGCC	1320
CTCTAAATAC	AAAAATAGCC	CTTCGGATAA	AATCCGAGGG	GCTAGAAACG	TTGTTAAATC	1380

310						
AACGGCCGAA	CTTTGAATT	TCATGGTCG	GGATAAAAATA	GTTCACTGAA	CTATTTTATT	1440
TTTTAAGGTT	ATCATAATAT	CAAATAGTTC	AATTAAATAC	GCTAAATTAC	TAATATACTT	1500
TTTACCTTT	TCATTCTAAA	ATGTAAGTA	CAAACAATTA	CAATATACTA	GAGGGGGAGT	1560
AAAAAAGGTA	TTAAATCGAT	GAGTTCAGCA	GGCAAGAAAA	TAGCACCTTT	ACGGGTGCTA	1620
TTTTTAATT	AACGCCACGT	TAACTTTGA	TTGATGAATT	TTATTGTTG	GCACTTCTT	1680
CATTTCACGG	AAACATCGA	TGAAATTCTT	TCCAACATTA	TTTTGGAGT	TAAC TG CATT	1740
TATTTTGTA	TTAATAACTT	TTTTAGTATC	GAAAGAATGG	TTTAAGAAAT	CCATAACTAA	1800
CTCTCCTTTC	TCATCCTGTA	ATCAAGATTT	TTATCAATGT	CAAATAGTA	TTTTCTATCA	1860
ATCCAAATTG	GTCCTTCTCC	TTAGAAATA	GCAAGTACAT	CTACCGGACC	TCCTACTGTT	1920
TCAAGAGTGT	TGACAATTTT	TCTCTTAAAT	GAAGTTAATT	CAATAATGT	TTAGCTGTA	1980
CTCGCCATT	CATTAAGTGG	TTGCATTCCA	ATAAGGTCTA	TTATAGGATT	TATATAATAT	2040
TTTTGCTGTA	TAGATGATAT	ATTTCAAAAT	ATATTCTCAA	TTTCATCACC	CAATCCATT	2100
TTCTCCATAA	CTGATGATAC	TTGCTCTGCG	ATATATACAT	TTAAGTTAGG	ATCTATACCA	2160
TTCATAATCG	TCTCAACCAT	CTCTGACTGT	GCAAAAGGGA	TTATATGACA	AGTTTTATGA	2220
TGATTATCA	CACTTCATT	AAATACTTTC	CAAATTAATC	GTTTAGAAAA	AATTCCATAT	2280
AATTCAATT	GTCTTATAGA	TGAAATATC	TCGTCTGTAC	CATAACCTGC	TATAACTAAT	2340
CCAGTTATGT	TTGTTGAGTC	ATATCCAATG	AAAATCGTT	TATATAAAGA	TTTAGCAATA	2400
ACTTCAACCT	CATCATCAGT	ATGAGGAAAG	GATTAAAAAA	CATCGTCTAC	AATGCTTTT	2460
ATTAACTCTA	ACTCAGCTTC	AAAAAATTCA	AAATTACTTT	CAGCTTCTAC	TTTTGAAATT	2520
TCTAAACTAA	ATTAGTTAT	AGCATTAAAT	AAAATTTTAT	TAAAATCATC	TAGAGTGATG	2580
GTTCACCAT	TAGAAACTCT	AAATCAGCT	GTTCCTTGCG	CTTCATAGGC	AATGCTGTCC	2640
AAAATACTTC	TTGTAACCTCT	GACAATATAA	TTCTTAATA	AATCCTCAAC	TTGTAAGATGT	2700
TTAAAGGAAA	TTAAAAATTC	TATTAAGCTTT	TCAACGTATT	GGGCAGTATT	ATCTAATAAA	2760
TCTGTGCCAA	TAGCTGCTT	AAACTCATTT	AAAATTACCT	CCCACGGAAT	TTCCATATAAC	2820
GAAGCGTTCC	CATATATCAT	GATCCCCACG	GAATGTTCTT	TTGATAAAAGT	GAATAATT	2880
CGGGCGCTAT	AAAAAACTTT	TGAATTCTTC	CCGCTGATA	AGGTTACAGC	GCTATCAGAA	2940
GCCAATACAA	CACCATT	TTAATATT	CCAATTCTG	CTGTCAAAAT	ATCACCTAAA	3000
CTTTCTAAAC	CTGCTCATGC	TCTAATGGTA	CAACAGCTAA	GGTCTTACCA	AGACTTGCCA	3060
ACACTTTAA	TACTGTATCA	AGTTGTGGGC	TTGTCTTCC	TGTTTCCATT	CTAGCGATAA	3120
CTGGCTGACT	AACACCGCTC	ATCTCCTCTA	GTTCCTCTG	ACTAATACCC	TTTTCAATT	3180

311

TAGCCTCGAT	AAGCTCACTC	ATGATAGCCA	CGCGCATATC	ACTTTCCAAA	ATTCCTCTT	3240
TGCTGAATAA	TTCAGCTCTT	ACATCTTCC	AGTTACTACC	AATAGCATT	TTTTCAT	3300
TCTAAACCTC	TTTCTTTAA	ATCTGCAAGT	TCACGTTAG	CTTGCTCAAT	CTCTCTTTG	3360
GGTGTTCCT	GTGTCCTTT	CATAAAATGA	TGCAGTAAA	CAAAACTACC	ATCCATCCAA	3420
GCAACAAATA	AAATTCTATC	TCTAAGTGGT	CTCAGCTCCC	AAATTCAGC	ATCTAAATGC	3480
TTAATATATG	GTCGCCTGC	GCGTGTCCA	TGTTGGCTTA	ACAACTCAAT	ATAATCATT	3540
ATTTTATTAA	GCTTAATTCT	GCTATCTTC	CCTTTTTAC	TGGTAAGCTC	TCGCATATAA	3600
TCAAAACAG	GCTCATTGCC	GTTTTATCC	TTGTAAAAT	AGATATTATG	CACTATTAAAC	3660
ACCTCTTCCT	ATAACAAATT	ATAACCTAA	AGTTATTGTT	TGTAAATACT	TTTAAGTTAT	3720
TAAAAATAAA	AGCACCTAGT	TTCCTAGATG	CTAGCACAAT	GACACGGATT	CGCACCGTGG	3780
CTACCTCTAT	CAAGGTGTAC	TCCTTCTATA	CTATCCCTG	TGCTTAGAA	TATTATACCA	3840
CACAATCAAC	TAGATACCTA	CCATCTCATG	ATATACCCC	ATTTGGGCA	AGGGTACAAC	3900
GCTAAATAC	AAATCAGAAT	AGATATTAA	CCACTTATT	AACTTATCAT	AAGCTGGTGA	3960
TTGACTGATA	ATAAATATCC	GCTGACAAGC	TCCGATAACA	TTCATGTGAT	TGTACACATA	4020
AACCTCTTT	ACAGCCTCTA	AAATGTCAGC	CTCACTGTGTT	TGTACCTAA	TATCTGTTAT	4080
CTGCTTGATA	GTTGCGTATT	TTTGATAAGC	TAGCATATCT	TGATTTTTAG	CAGCATCAA	4140
CATTTTACGC	TCAAGGACAC	TATACCTAGG	TTGTTCTTTA	TCTCGCATGA	AATACCACTT	4200
GAGCCATAAA	ATCTTTCTC	GGTGTATTAC	AGAAATACGC	TCAATTTCT	TCTTGTCAT	4260
TGCTACCTCC	AAATCATCA	ATTAACAAT	TCTAACCACT	CACTTTAGA	AATAGTTGCA	4320
TAGATCTTGT	TCGATGTATG	ATACAAAGGT	TCTAAATCTT	TTTCCACCT	AATATAGTT	4380
ATCTTATCCT	CATGAGTAGG	AAAGTATAGT	ATTTCCGTT	CATCCTCGTT	TAGGATACGA	4440
TTGCACCAAT	CATCAATAAT	AACTGGCACT	TCCCACTCAC	GCCATTTTT	AAGGTTTTCT	4500
AAAAGTTCAT	TATCACTAAA	TAGCTCGCCA	TCTATTTGGA	AAAATCCCC	TAAGTCATTG	4560
TTTCCTCAA	CAATAATAAA	CTCTGGCATA	TTTCTATTAC	TTAATAACTC	CTTGAGTTCT	4620
TGTAACCTCTT	TGATTTCTT	TAGATACTTC	CTCAATTCC	AACCTCAATT	CTTCAATCTG	4680
CCTTACTACT	CCAAAAATT	CATGGGTCTT	ATAAGATTGT	TCAAGTATAG	CCTTGCTGC	4740
TTGAGTTCTT	ATAAACGGGT	TGACCTTA	GTCCATCATA	ATATCATTGA	GTACAGAAC	4800
AGCGTTAGAT	GATGCTAAAT	AAAGCATTG	AGTTGTTTA	TCCATCATCT	CATCTGCTT	4860
TATCCTCAAT	GTCTTTAA	CCGCTGCAAC	TTTAGATAC	TTATGACCTG	TTGCGCGTGA	4920

312						
TACCCCTGCT	TTTGACATG	CTTGTCTAT	CGTTGGCTCG	GTAAGCATGG	CATCTATGAA	4980
TTTAATTG	TTGGACGTAA	GGTTATCATT	TTCATTCCT	GCCATCTATT	ACCTCCTCAT	5040
TATCAAATA	AAGGGTTGCC	CCTTTATTTC	CCTATGCTAG	ATAATTCTGC	AATTCTGCAT	5100
CCATTGCCTC	TGAATTGCC	TCAACAATCA	TTTCATGCTG	TACTAAATCA	ATCTTATCTC	5160
CGTTAATAAG	TAAACCACCG	TGGAAATAAT	CAATTTTCT	ATCAAGGAAA	TGTACTAGCT	5220
TTTCAAGCG	TTGCTGTTGG	CTGAATTGCT	CCATGTCAAT	TTCGATATAA	GCAAGGGTAG	5280
TATCATTATC	CATAATATCT	TCTAATTTC	TAAGAGCTAG	AGGTTTATT	TTATATT	5340
CTAGGTATTC	TCTCATTCT	GCCACTGTTA	ATTTGATACT	AGATAATAAA	CTTAGTTCA	5400
CTGCATC	TGCTGTAATA	GGCTCTTCTT	TTGATTCATG	TTTGCTAGT	TCAGCATTTT	5460
TCTCTTTTC	TAGTGCTGA	TACAATAGCT	GAGCAGTATT	TTGGGAATAG	TTTCGCCCT	5520
CTTTTTATA	TTTTAAAAGT	TCTTGCTCTG	CATACACTTT	CCCGATAATC	ACTTCCTTAT	5580
AAACTAATTG	CCCATCTG	GCTTTAGCT	TAATACTCCC	ATGCTCTGGA	ATTTCAATAT	5640
ACTTAATTAT	ACCATT	GAGTATAAAA	CAAAGCCTT	CTCCATCATT	TTTAATAATT	5700
TATCATC	CTTTCAGTC	ATGCTTTCT	CCTTATTTC	ATTTTATTAT	AATCTGAATA	5760
CCCCTAGTCT	ATTTATTCA	CTAGGTTTTT	AGGGTCGTA	TGCTAAAATA	CTACCTTTT	5820
TGTGTAC	CTTCAATTG	GACT	TTTAGTT			5857

(2) INFORMATION FOR SEQ ID NO: 29:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 10254 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 29:

AAAATGATAG	CAGGAGAGTT	TTCCCGTCCA	TCAGACCCAG	AACTGAGAGC	CTTAGCTCAG	60
GCTTCTCGCC	AAAAACAGGC	CGCCTTTAAC	AAGGAAGAGA	ACCCCTTGAA	GGGAGGCCGAA	120
ATCATCAAGA	CTTGGTTGC	CTCAACCGGG	AAAAATCTTT	ACATCAACAC	TCGCTTGATG	180
GTGGACTACG	GTGTCAACAT	CCATCTAGGG	AAAAATT	ATTCTAATTG	GAACTTGACC	240
ATGCTGGATA	TCTGTCCCAT	TCGTATCGGG	GACAATGCTA	TGATTGGTCC	TAATTGTCAG	300
TTTTGACAC	CCCTCCATCC	ACTAGATCCA	CAGGAACGCA	ATTCAGGTAT	CGAGTACGGA	360
AAGCCTATCA	CAATCGGAGA	TAATTCTGG	ACTGGTGGTG	CGTCATTGT	CCTTCCTGGA	420
GTGACACTGG	GAAATAATGT	CGTTGCAGGA	GCAGGGCAG	TAATTACCAA	ATCTTTGGC	480

GACAACGTTG	TCCTAGCTGG	CAATCCTGCG	CCCGTGATTA	AGGAAATACC	TGTTAAATAG	540
AAGTAAAAG	GAACAGCTGG	GGTTGTTCT	TTTTGTAGG	TTTCATCATT	TTTTACCCAG	600
TTCACATT	A CCTACTCTAT	CTCTTAGCAA	GTCTGTTCA	TTAAGCAGT	TCAAAGCATC	660
TCGTAAGTGG	GATGTTTTC	TCCTCAGTTC	ATCAGCTTC	TCCTTGACAC	TCGGTCAGAT	720
TTTGATACAA	TAGTACAAA	TTAGAGGAGG	CAGGCTATGA	TTCAGAAACA	TGCGATTCT	780
ATTTTAGAGT	TTGATGACAA	TCCTCAGGCG	GTATCATGC	CCAATCACGA	GGGGCTGGAC	840
TTGCAGTTGC	CAAAGAAGTG	TGTTTATGCA	TTTTTAGGTG	AGGAGATTGA	CCGCTATGCG	900
AGGGAAAGTAG	GGCGAAGCTG	TGTTGGCAA	TTTGTGTTCTG	CCACCAAGAC	CTATCCAGTT	960
TATGTCGTGA	ACTACAAGGA	CGAGGAGGTC	TGTCCTGGCTC	AGGCTCCTGT	TGGCTCCGCT	1020
CCAGCAGCCC	AGTTATGGA	TTGGTTGATT	GGCTATGGTG	TGGAGCAGAT	TATCTCTACT	1080
GGGACCTGTG	G TGTCCTAGC	TGATATAGAG	GAAAATGCCT	TTCTAGTCCC	TGTTCGCGCT	1140
CTGCGAGATG	AAGGAGCCAG	TTACCACTAT	GTGGCACCTT	GTGTTATAT	GGAAATGCAG	1200
CCAGAGGCTA	TTGCTGCTAT	TGAGGAAGTT	TTGGAAGACA	GAGGGATTCC	TTATGAAGAA	1260
GTCATGACCT	GGACGACAGA	CGGTTTTAC	CGAGAACCG	CTGAAAAGGT	GGCTTATCGT	1320
AAGGAAGAAG	GCTGTCGTG	TGTTGGAGATG	GAGTGTCTG	CTCTTGCAGC	AGTAGCTCAA	1380
TTGCCTGGGG	TTCTCTGGGG	TGAATTGTTG	TTCACACCGAG	ATTCTCTAGC	GGACTTGGAC	1440
CAGTACGACA	GTCGTGACTG	GGCTCGCAA	CCCTTTAATA	AGGCCTAGA	ACTGAGTTA	1500
GCAAGTGTTC	ACCACCTTA	GTTGTACTGG	CAAAGGATT	CTTTTATCAT	AAAATGTCTA	1560
GCTCATACTT	TTCAAAAATA	TGTTAAACG	AGGTACACCTT	CCTCTTGTCC	TAGGCATGTT	1620
GAGGTTGGGA	AAAATTTA	AAATCAGAAA	AACGTATCAT	ATCAGGTGAT	AAAAACTTG	1680
ACACTATGCG	TTTTATGTCG	ATAAGATT	GAGTGAGATG	AAATGATACT	CTTCGAAAAT	1740
CTCTTCAAAC	CAGGTCAAGT	TCACCTTGCC	GTAGGTATAT	GTACTGACT	TCGTCAGTCT	1800
TATCCGGCAA	CCTCAAAACG	GTGTTTGAG	CTGACTTCGT	CAGTTCTATT	TGCAACCTCA	1860
AAACAGTGTT	TTGACCAACC	TGTGACTAGC	TTTCTAATCG	ATGCCTTGGT	TTTCATTGCC	1920
TATAATCAA	AAGAGAAATT	TTCTCCCTGAA	AAGCATATAG	AGTAGCTGGC	GTTAAAAGCT	1980
CCTGTCTTGC	TTTTTGACC	TATAGTCACA	TCTATCAAGT	ATTGTTCTTG	CCTAAGCTAT	2040
CAATAAAAAG	GTGGCATT	TTAGGCTTGG	TGTTAGTAGA	TTTTGCCTTA	TCCTATCTAA	2100
GTCATTTCGA	ACTTTTATG	GTACAATGGA	AACATGTTAT	TCAAATTATC	TAAGGAAAAA	2160
ATAGAGCTAG	GCTTATCTCG	TTTATGCCA	GCCCGTCGTA	TTTTTTGAG	TTTGCCTTG	2220

314			
GTCATTTAC TAGGCTCTCT	TCTTTGAGC TTGCCCTTG	TCCAAGTTGA AAGCTCACGA	2280
GCGACTTATT TTGATCATCT	TTTCACTGCT GTCTCTGCAG	TCTGTGTGAC GGGCTCTCA	2340
ACCCTTCCAG TAGCTCACAC	CTATAATATC TGGGGTCAAA	TAATCTGTT GCTCTTGATT	2400
CAGATCGGTG GTCTAGGGCT	CATGACCTTT ATTGGGGTTT	TCTATATCCA GAGCAAGCAA	2460
AAGCTAGTC TTCGTAGCCG	TGCAACTATT CAGGATAGTT	TTAGTTATGG AGAAACTCGA	2520
TCTTTGAGAA AGTTTGTCTA	TTCTATTTTT CTCACGACCT	TTTGGTTGA GAGCTGGGA	2580
GCTATTTGC TTAGTTTCG	CCTTATTCCCT CAACTGGCT	GGGGACGTGG TCTTTTTAGT	2640
TCCATTTTC TAGCGATCTC	AGCCTCTGT AATGCCGGTT	TTGATAATT AGGGAGCAC	2700
AGTTTATTTG CTTTCAGAC	CGATTTACTG GTCAATCTGG	TGATTGCAGG CTTGATTATT	2760
ACAGGGCGGCC TTGGTTTTAT	GGTCTGGTTT GATTGGCTG	GTCATGTAGG AAGAAAGAAA	2820
AAAGGACGTC TGCACTTCA	TACGAAGCTT GTACTATTAT	TGACTATAGG TTTGTTGTTA	2880
TTTGGAACAG CAACTACTCT	CTTTCTTGAG TGGAACAAATG	CTGGAACGAT TGGCAATCTC	2940
CCTGTTGCCG ATAAGGTTTT	AGTTAGCTTT TTTCAAACAG	TGACGATGCG AACAGCTGGC	3000
TTTTCTACGA TAGATTATAC	TCAGGCTCAT CCTGTGACTC	TTTGATTTA TATCTTACAG	3060
ATGTTTCTAG GTGGGGCACC	TGGAGGAACA GCTGGGGAC	TCAAGATTAC GACATTTTT	3120
GTCCTCTTGG TCTTGCACG	AACTGAGCTT CTAGGCTTGC	CTCATGCCA TGTTGCGAGA	3180
CGAACGATCG CGCCGCGAAC	GGTTCAAAAA TCCTTAGTG	TCTTTATTAT CTTTTGATG	3240
AGCTTCTTGA TAGGATTGAT	TCTGCTAGGG ATAACAGCCA	AAGGCAATCC TCCCTTATC	3300
CACCTCGTAT TTGAAACCAT	TTCAAGCTTT AGTACAGTTG	GTGTAACGGC AAATCTGACT	3360
CCTGACCTTG GGAAATTGGC	TCTCAGTGTT ATCATGCCA	TTATGTTAT GGGACGAATT	3420
GGTCCCTTGA CCTTGTGTTGT	TAGCTGGCA GATTACCATC	CAGAAAAGAA AGATATGATT	3480
CACTATATGA AAGCAGATAT	TAGTATTGTT TAAGAAAGGA	AAGAGCATGT CAGATCGTAC	3540
GATTGGAATT TTGGGCTTGG	GAATTTTG GAGCAGTGT	CTAGCTGCC TAGCCAAGCA	3600
GGATATGAAT ATTATCGCTA	TTGATGACCA CGCAGAGCGC	ATCAATCAGT TTGAGCCAGT	3660
TTTGGCCGCT GGAGTGATTG	GTGACATCAC AGATGAAGAA	TTATTGAGAT CAGCAGGGAT	3720
TGATACCTGC GATACCGTTG	TAGTCGCGAC AGGTGAAAAT	CTGGAGTCGA GTGTGCTTGC	3780
CGTTATGCAC TGTAAAGAGTT	TGGGGGTACC GACTGTTATT	GCTAAGGTCA AAAGTCAGAC	3840
CGCTAAGAAA GTGCTAGAAA	AGATTGGAGC TGACTCGTTT	ATCTCGCCAG AGTATGAAAT	3900
GGGGCAGTCT CTAGCACAGA	CCATTCTTTT CCATAATAGT	GTTGATGTCT TTCAGTTGGA	3960
TAAAAATGTG TCTATCGTGG	AGATGAAAAT TCCTCAGTCT	TGGGCAGGTC AAAGTCTGAG	4020

TAAATTAGAC	CTCCGTGGCA	AATACAATCT	GAATATTTG	GGTTTCCGAG	AGCAGGAAAA	4080
TTCCCCATTG	GATGTTGAAT	TTGGACCAGA	TGACCTCTTG	AAAGCAGATA	CCTATATTTT	4140
GGCAGTCATC	AACAACCACT	ATTGGATAC	CCTAGTAGCA	TTGAATTCTGT	AAAGAGGGAT	4200
GACCCCTCTT	TTTGATGCC	TAAGATGGCA	AATAGAGACA	GAAGCCCCTT	GTCTTCTAGT	4260
AAAAGTTCTT	CAAAGGCTGG	ACTTTATGGT	AAAATAGAAA	GAAGTGCACAA	GAGAGAGTAA	4320
TAATCAATGA	AAATCAAAGA	TCAAACCTAGG	AAACTAGCTA	CGGGCTGCTC	AAAACACTGT	4380
TTTGAGGTTG	CAGATAGAAC	TGACGAAGTC	AGTAACATCT	ATACGGCAAG	GCGACGTTGA	4440
CCCGGTTTGA	AGAGATTTTC	GAAGAGTATA	AGAAAAAAATC	AGTCCCCTAA	AGGAGTAGAT	4500
TATGAAGTTA	TTGCTCTATCG	CAATTTCTAG	CTATAATGCA	GCAGCCTATC	TTCATTACTG	4560
TGTGGAGTCG	CTAGTGATTG	GTGGTGAGCA	AGTTGGGATT	TTGATTATCA	ATGACGGGTC	4620
TCAGGATCAG	ACTCAGGAAA	TCGCTGAGTG	TTTAGCTAGC	AACTATCCTA	ATATCGTTAG	4680
AGCCATCTAT	CAGGAAAATA	AATGCCATGG	CGGTGCGGTC	AATCGTGGCT	TGGTAGAGGC	4740
TTCTGGCGC	TATTTTAAAG	TAGTTGACAG	TGATGACTGG	GTGGATCCTC	GTGCCTACTT	4800
GAAAATTCTT	GAAACCTTGC	AGGAACCTGA	GAGCAAAGGT	CAAGAGGTGG	ATGTCTTTGT	4860
GACCAATTCTT	GTCTATGAAA	AGGAAGGGCA	GTCTCGTAAG	AAGAGTATCA	GTTACGATTC	4920
AGTCTTGCT	GTTCCGCAGA	TTTTGGCTG	GGACCAGTC	GGAAATTCT	CCAAAGGCCA	4980
GTATACCTATC	ATCCACTCGC	TGATTTATCG	GACAGATTTG	TTGGGTGCTA	GCCAGTTCTA	5040
ACTGCCTGAA	CATACTTTT	ATGTCGATAA	TCTCTTTGTC	TTTACGCC	TTCAGCAGGT	5100
CAAGACCATG	TACTATCTGC	CTGTCGATTT	CTATCGTT	TTGATTGGGC	GTGAGGACCA	5160
GTCTGTCAAT	GAGCAAGTGA	TGATTAAGTG	CATTGACCAG	CAACTCAAGG	TCAATCGACT	5220
CTTGATAGAC	CAACTTGATT	TGTCCCAAGT	GAGTCATCCC	AAAATGCGAG	AATATCTGCT	5280
GAATCATATT	GAACTCACGA	CGGTGATTTC	CAGTACCCCTG	CTCAACCGAT	CTGGAACAGC	5340
GGAGCATCTG	GCAAAAAAAC	GCCAATTGTG	GACCTATATT	CAGCAGAAAA	ATCCAGAAGT	5400
CTTTCAGGCT	ATTCGTAAGA	CCATGTTGAG	CCGTTTGACC	AAACATTCTG	TCTTGCCAGA	5460
TCGCAAACCTG	TCCAATGTCG	TCTATCAAAT	CACCAAATCT	GTTCATGGAT	TTAATTAATA	5520
TAAGTGTGTTT	ATAAGAGGGA	TTTAAGAAAA	ATTTTAACCT	TTTCTTAGTC	CTTTTTAATT	5580
TCAGGAGATT	ATACTAGAGT	CATCAAATAA	AGAAAGACTC	TAAGGAGAAT	CCTATGAAAT	5640
TCAATCCAAA	TCAAAGATAT	ACTCGTTGGT	CTATTCGCCG	TCTCAGTGTG	GGTGTGCGCT	5700
CAGTTGTTGT	GGCTAGTGGC	TTCTTTGTCC	TAGTTGGTCA	GCCAAGTTCT	GTACGTGCCG	5760

316	
ATGGGCTCAA TCCAACCCCA GGTCAAGTCT TACCTGAAGA GACATCGGG ACGAAAGAGG	5820
GTGACTTATC AGAAAAACCA GGAGACACCG TTCTCACTCA AGCGAACCT GAGGGCGTTA	5880
CTGGAAATAC GAATTCACTT CCGACACCTA CAGAAAGAAC TGAAGTGAGC GAGGAAACAA	5940
GCCCTCTAG TCTGGATACA CTTTTGAAA AAGATGAAGA AGCTAAAAAA AATCCAGAGC	6000
TAACAGATGT CTTAAAAGAA ACTGTAGATA CAGCTGATGT GGATGGGACA CAAGCAAGTC	6060
CAGCAGAAAC TACTCCTGAA CAAGTAAAAG GTGGAGTGAA AGAAAATACA AAAGACAGCA	6120
TCGATGTTCC TGCTGCTTAT CTTGAAAAAG CTGAAGGGAA AGGTCCCTTC ACTGCCGGTG	6180
TAAACCAAGT AATTCTTAT GAACTATTGCT CGTGTGATGG TATGTTAACT CGTCTATTAC	6240
TAAAAGCTTC GGATAATGCT CCTTGGTCTG ACAATGGTAC TGCTAAAAAT CCTGCTTTAC	6300
CTCCTCTTGA AGGATTAACA AAAGGGAAAT ACTTCTATGA AGTAGACTTA AATGGCAATA	6360
CTGTTGGTAA ACAAGGTCAA GCTTTAATTG ATCAACTTCG CGCTAATGGT ACTCAAACTT	6420
ATAAAAGCTAC TGTTAAAGTT TACGGAAATA AAGACGGTAA AGCTGACTTG ACTAATCTAG	6480
TTGCTACTAA AAATGTAGAC ATCAACATCA ATGGATTAGT TGCTAAAGAA ACAGTTCAAA	6540
AAGCCGTTGC AGACAACGTT AAAGACAGTA TCGATGTTCC AGCAGCCTAC CTAGAAAAAG	6600
CCAAGGGTGA AGGTCCATTC ACAGCAGGTG TCAACCATGT GATTCCATAC GAACTCTTCG	6660
CAGGTGATGG CATGTTGACT CGTCTCTTGC TCAAGGCATC TGACAAGGCA CCATGGTCAC	6720
ATAACGGCGA CGCTAAAAAC CCAGCCCTAT CTCCACTAGG CGAAAACGTG AAGACCAAAG	6780
GTCAATACTT CTATCAAGTA GCCTTGGACG GAAATGTAGC TGCGAAAGAA AAACAAGCGC	6840
TCATTGACCA GTTCCGAGCA AAYGGTACTC AAACTTACAG CGCTACAGTC AATGTCTATG	6900
GTAACAAAGA CGGTAAACCA GACTTGGACA ACATCGTAGC AACTAAAAAA GTCACTATTA	6960
ACATAAACGG TTTAATTCTTAAAGAAACAG TTCAAAAGC CGTTGCAGAC AACGTTAAAG	7020
ACAGTATCGA TGTTCCAGCA GCCTACCTAG AAAAGCCAA GGGTGAAGGT CCATTACAG	7080
CAGGTGTCAA CCATGTGATT CCATACGAAC TCTTCGCAGG TGATGGTATG TTGACTCGTC	7140
TCTTGCTCAA GGCATCTGAC AAGGCACCAT GGTCAGATAA CGGTGACCGCT AAAAACCCAG	7200
CCCTATCTCC ACTAGGTGAA AACGTGAAGA CCAAAGGTCA ATACTTCTAT CAATTAGCCT	7260
TGGACGGAAA TGTAGCTGGC AAAGAAAAAC AAGCGCTCAT TGACCAGTTC CGAGCAAACG	7320
GTACTCAAAC TTACAGCGCT ACAGTCAATG TCTATGGTAA CAAAGACGGT AAACCAGACT	7380
TGGACAAACAT CGTAGCAACT AAAAGTCA CTATTAACAT AAACGGTTTA ATTTCTAAAG	7440
AAACAGTTCA AAAAGCCGTT GCAGACAACG TTAAGGACAG TATCGATGTT CCAGCAGCCT	7500
ACCTAGAAAA GGCCAAGGGT GAAGGTCCAT TCACAGCAGG TGTCACCAGT GTGATTCCAT	7560

317

ACGAACCTCTT	CGCAGGTGAT	GGCATGTTGA	CTCGTCTCTT	GCTCAAGGCA	TCTGACAAGG	7620
CACCATGGTC	AGATAACGGC	GACGCTAAAA	ACCCAGCTCT	ATCTCCACTA	GGTGAACCG	7680
TGAAGACCAA	AGGTCATAAC	TTCTATCAAG	TAGCCTTGGA	CGGAAATGTA	GCTGGCAAAG	7740
AAAAACAAGC	GCTCATTGAC	CAGTTCCGAG	CAAACGGTAC	TCAAACATTAC	AGCGCTACAG	7800
TCAATGTCTA	TGGTAACAAA	GACGGTAAAC	CAGACTTGGA	CAACATCGTA	GCAACTAAAA	7860
AAGTCACTAT	TAAGATAAAAT	GTAAAGAAA	CATCAGACAC	AGCAAATGGT	TCATTATCAC	7920
CTTCTAACTC	TGGTCTGGC	GTGACTCCGA	TGAATCACAA	TCATGCTACA	GGTACTACAG	7980
ATAGCATGCC	TGCTGACACC	ATGACAAGTT	CTACCAACAC	GATGGCAGGT	GAAAACATGG	8040
CTGCTCTGC	TAACAAGATG	TCTGATACGA	TGATGTCAGA	GGATAAGCT	ATGCTACCAA	8100
ATACTGGTGA	GACTCAAACA	TCAATGGCAA	GTATTGGTT	CCTTGGGCTT	GCGCTTGCAG	8160
GTTTACTCGG	TGGTCTAGGT	TTGAAAACA	AAAAAGAAGA	AAACTAATCA	GCTAAGGAAA	8220
TAAATGATGG	ATAGTGGGCT	GACTAAGATT	AGTTTAACAA	CTCAATCAGC	AATCAGGACT	8280
TTCTTCAAT	AGCAGATTAA	AATCATCGTA	AAACAATAAA	AATAGTGTAA	TACTTAAAGC	8340
AGTATAGCAC	TGTTTTATC	AAAGGAGAGA	CAGATGGAA	AGACAATTTT	ACTCGTTGAC	8400
GACGAGGTAG	AAATCACAGA	TATTCATCAG	AGATACTTAA	TTCAGGCAGG	TTATCAGGTC	8460
TTGGTAGCCC	ATGATGGACT	GGAAAGCGCTA	GAGCTGTTCA	AGAAAAAAC	GATTGATTG	8520
ATTATCACAG	ATGTCATGAT	GCCTCGGATG	GATGGTTATG	ATTTAATCAG	TGAGGTTCAA	8580
TACTTATCAC	CAGAGCAGCC	TTTCCTATTT	ATTACTGCTA	AGACCAGTGA	ACAGGACAAG	8640
ATTTACGGCC	TGAGCTGGG	AGCAGATGAT	TTTATTGCTA	AGCCTTTAG	CCCACGTGAG	8700
CTGGTTTGC	GTGTCCACAA	TATTTGCGC	CCCTTCATC	GTGGGGCGA	AACAGAGCTG	8760
ATTTCCCTTG	GCAACTAAA	AATGAATCAT	AGTAGTCATG	AAGTTCAAAT	AGGAGAAGAA	8820
ATGCTGGATT	TAACTGTTAA	ATCATTGAA	TTGCTGTGGA	TTTTAGCTAG	TAATCCAGAG	8880
CGAGTTTCT	CCAAGACAGA	CCTCTATGAA	AAGATCTGGA	AAGAAGACTA	CGTGGATGAC	8940
ACCAATACCT	TGAATGTGCA	TATCCATGCT	CTTCGACAGG	AGCTGGCAA	ATATAGTAGT	9000
GACCAAACCTC	CCACTATTAA	GACAGTTGG	GGGTTGGGAT	ATAAGATAGA	GAAACCGAGA	9060
GGACAAACAT	GAAACTAAAA	AGTTATATTT	TGGTTGGATA	TATTATTCA	ACCCTCTTAA	9120
CCATTTGGGT	TGTTTTTGG	GCTGTTCAA	AAATGCTGAT	TGCGAAAGGC	GAGATTTACT	9180
TTTTGCTTGG	GATGACCATC	GTTGCCAGCC	TTGTCGGTGC	TGGGATTAGT	CTCTTCTCC	9240
TATTGCCAGT	CTTTACGTG	TTGGGAAAC	TCAAGGAGCA	TGCCAAGCGG	GTAGCGGCCA	9300

318

AGGATTTCC	TTCAAATTG	GAGGTTCAAG	GTCCTGTAGA	ATTCAGCAA	TTAGGGCAA	9360
CTTTTAATGA	GATGTCCCCT	GATTTGCAGG	TAAGCTTGA	TTCCCTGGAA	GAAAGCGAAC	9420
GAGAAAAGGG	CTTGATGATT	GCCAGTTGT	CGCATGATAT	TAAGACTCCT	ATCACTTCGA	9480
TCCAAGCGAC	GGTAGAAGGG	ATTTGGATG	GGATTATCAA	GGAGTCGGAG	CAAGCTCATT	9540
ATCTAGCAAC	CATTGGACGC	CAGACGGAGA	GGCTCAATAA	ACTGGTTGAG	GAGTTGAATT	9600
TTTGACCCCT	AAACACAGCT	AGAAAATCAGG	TGGAAACTAC	CAGTAAAGAC	AGTATTTTC	9660
TGGACAAAGCT	CTTAATTGAG	TGCATGAGTG	AATTCAGTT	TTTGATTGAG	CAGGAGAGAA	9720
GAGATGTCGA	CTTGCAAGGT	ATCCCAGAGT	CTGCCCGGAT	TGAGGGAGAT	TATGCTAACG	9780
TTTCTCGTAT	CTTGGTGAAT	CTGGTCGATA	ACGCTTTAA	ATATTCTGCT	CCAGGAACCA	9840
AGCTGGAAGT	GGTGGCTAAG	CTGGAGAAGG	ACCAGCTTTC	AATCAGTGTG	ACCGATGAAG	9900
GGCAGGGTAT	TGCCCCAGAG	GATTGGAAA	ATATTTCAA	ACGCCCTTAT	CGTGTGAA	9960
CTTCGCGTAA	CATGAAGACA	GGTGGTCATG	GATTAGGACT	TGCGATTGCG	CGTGAATTGG	10020
CCCATCAATT	GGGGGGGAA	ATCACAGTCA	GCAGCCAGTA	CGGTCTAGGA	AGTACCTTA	10080
CCCTCGTTCT	CAACCTCT	GGTAGTGAAA	ATAAAGCCTA	AAACCCCTTT	ACAAATCCAG	10140
CTATTCAATGG	TAGAATAGAT	TTTGTGTGAA	ATATCAGCAG	GAAAGCATGA	AGCTCGTCAA	10200
CAGGTGTCTT	ATGACAAGTA	ACCTGGCTG	TTTAGGCAGA	GGGCATCTGC	ACGG	10254

(2) INFORMATION FOR SEQ ID NO: 30:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 9769 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 30:

CCGGCGACTA	TCGATAACAC	TTGACTTGGT	AGCCCCACAT	TTGGACAAC	GCATCCTTTC	60
CCTCCTTATC	GTTCCTTTT	CATTATACCA	TTTTTAAGC	GATTCCAAA	ACAATTCTTC	120
TTTTGCTTG	ACAAGTTTT	TGTTTGTTG	TATTATTTAA	TTAAGACAAC	AAGGTAAAAG	180
AAAGGAGACT	AAGATGTCCT	GGACATTGTA	CAACAAAAAA	CCCATCTATT	TACAGATTAT	240
GGAGAAAATC	AAGCTTCAGA	TTGTTTCCCA	TACACTGGAA	CCCAATCAAC	AACTTCCAAC	300
CGTGAGGAGC	TAGCTAGCGA	GGCTGGGTGTC	AATCCAATA	CCATCCAAAG	AGCCTTATCA	360
GACCTTGAAC	GAGAAGGATT	TGTCTACAGC	AAGCGAACAA	CTGGACGATT	TGTGACTAAG	420
GATAAGGAGC	TAATCGCCCA	GTCACGCAA	CAATTATCAG	AAGAAGAATT	GGAAACACTTC	480

GTTCCTCCA TGACCCATTT TGGCTATGAA AAAGAAGAAC TACCAGGCGT AGTCAGTGAT	540
TATATTAAG GAGTTAACG CTATGTCATT ACTAGTATTT GAAAATGTAT CCAAATCATA	600
TGGAGCAACA CCAGCCCTTG AAAATGTTTC TCTTGACATT CCAGCTGGAA AAATTGTCGG	660
CCTTCTTGGG CCAAACGGCT CAGGAAAAAC AACCTGATT AACTAATTA ATGGCCTCTT	720
ACAACCAGAT CAAGGACGTG TCCTCATCAA CGACATGGAC CCAAGCCCAG CAACCAAGGC	780
CGTTGTAGCT TATTGCTTG ATACGACCTA TCTCAATGAG CAAATGAAGG TCAAAGAACG	840
CCTAACCTAC TTCAAGACCT TCTATAAAGA TTGTCAGATC TTGAACGCGC CCATCATCTA	900
CTTGCAGACC TGGGCATTGA TGAAAATAGT CGTCTCAAGA AACTATCAAAGGAAACAAA	960
GAAAAGGTTTC AACTGATTTT CGTTATGAGC CGTGATGCTC GTCTCTATGT TTTGGACGAA	1020
CCCATTGGTG GGGTGGATCC AGCAGCCCGT GCTTATATCC TCAATACCAT TATCAACAAAC	1080
TACTCACCAA CTTCTACCGT TTTGATTTCT ACCCACTTGA TTTCTGATAT CGAGCCAATC	1140
TTGGATGAAA TTGTCCTCCT AAAAGACGGA AAAGTCGTCC GTCAAGGAAA TGTAGATGAT	1200
ATTCGCTACG AGTCAGGTGA ATCCATTGAC CAACTCTTCC GTCAGAATTT AAGGCCTAAG	1260
CAAAGGAGAT TATTATGTT TTGGAATTAA GTTCGCTACG AATTAAAAA TGTTAACAAAG	1320
TGGTATTTAG CCCTCTACGC AGCCGTGCTA GTCCCTTCTG CCCTCATCGG AATACAGACA	1380
CAAGGCTTTA AAAATCTACC TTACCAAGAA AGTCAGGCTA CTATGCTACT TTTCTAGCT	1440
ACAGTCTTTCG GTGGCTTGAT CCTACACTT GGGAATTCAA CCATTTCTT GATTATTAAG	1500
CGCTTCAAAG GTAGTGTCTA CGACCGACAA GGCTATCTGA CTTTGACCTT GCCAGTTCT	1560
GAACACCATA TCATCACAGC CAAACTAATC GGTGCCTTTA TCTGGTCATT GATTAGCACC	1620
GCTGTATTGG CTCTAAGTGC TGTATTATTCTGGCTTTAA CAGCTCCAGA ATGGATTCC	1680
CTTTCTTATG TGATTACATT TGTAGAAACA CATCTCCCTC AGATCTTTCT TACAGGTATA	1740
TCCTTCCTAC TAAATACTAT TTCAAGAAC CTCTGCATCT ACCTGGCTAT TTCCATTGGA	1800
CAGCTTTCA ATGAATACCG TACAGCACTC GCTGTTGCAG TCTACATTGG TATCCAAATC	1860
GTCATTGGAT TTATTGAACCTTCTCAAT CTTAGTTCTA ATTTCTATGT CAATTCACTG	1920
GTAGGACTCA ATGACCACATT CTATATGGGA CCAGGTATAG CCATTGTTGA AGAACTCATA	1980
TTCATAGCTA TCTTTATCT CGGAACCTAC TACATCTTGA GAAATAAGGT TAATTGCTT	2040
TAAATAATTT TTACCTAGAT ATGTAACATA CTCTAGAAC AAAAGAGACC AGGCAAAAG	2100
TCTTTAAAT TAGAAAACGC ATAGTATCAG GTGTTGAATA TGTACTGCC CCCCCAAAGTT	2160
AGATTTTTTC TGTCTAACTT TTGGGGCAG TTCATAAGAA CCTTGTTAAT ATGCGTTTT	2220

320	
TGTGAGCTGA CTTATTTCTT TTCACTATAT CGCAAAATGA AATAAGAACG GAACGATGGG	2280
ATTTTGAAT TCAAATCAAT TTATAAGAAT GTTTTAGAAG TAATATTATC CTATTCCAGA	2340
TTCAGTTAC TATACAATTG ACTTTCAAG CAACCTGTT ACATAATGTG TACATAATT	2400
GGTTCCGTAT TCCACCCCTT TCACCTTTAA AACCTCGCT TTGCAAGGC TCTTCTATT	2460
ATAAGATAAG GCACGTTAA AGGTTTCCA AATCCCTAAA TCATCCGTT GAAGAACGAG	2520
ACTAGCATAAC ATGCGTCGA TAAATCCTGT TGCTACCACC GCAAAATCA CTGTAATAGC	2580
AAGTGAAATC CATGCTTCTG CTCCCCCGC ATAGTCATTA ATCGTTGAA ACGGCATAAA	2640
GAAGGTCGAA ATAAAGGAA TATAAGAACC AATCTCAAG AGGAGATTGT CACCAGCTGC	2700
ACCTAGAGCT GTCACTCCAA AAAAACACC CATAATCAAATCATAAG GCGACAAGGC	2760
TTTCCCTGAG TCCTCAGGAC GAGAAACCAT AGATCCTAGG AAGGCTGCCA AGACTACGTA	2820
CATGAAAAGA CTGATCAAAA TAAAGAGCAA GGTATTCACT GAGATAGCAT CTCCCAAGTG	2880
ATCCAAAATA CCAGACTGAG CCAAGAATGG CAAATCTTTA AAGAGCAAA CGGCAGCCAG	2940
ACCACCTACA ACATAGATCC CAATATGCGT TAAAATCACT AGAAACAGAG CCATCATCCG	3000
CGCATAGAAA TAGTGAATTG CCCTTATGCT AGAAAAAACG ACTTCCATAA TTTTGGTGC	3060
TTTTTCACTG GCAACTTCCT GAGCTGTTAC ACCCGCATAG GTAATCAGAA TCATATAAAG	3120
AAAGAATCCT AAGGCACCTG CTGCAATTGT TTGAATAAAC TTTTATTTT CTTGGCTTC	3180
ATCAATCTTT TCTGTGAATT GAATTGTCTG CGCTAAGCGT TTTCTGCT CTTGAGACAA	3240
GGAAAGCAGTT GAAGCATTAA GCTGATTTTG CAGTTCATG AGTGTACCTG TAACCTCAAA	3300
TTTAATTCCA TTTCAAGCG ATGTTTCGCC ATGATAAACT GCCTTTAGAA CACTATCTTC	3360
TTGATCAATG GTCAAATAAC CTTTTAATTT TTCTCTTTA ATTGCTTCTT TGGCACTTGC	3420
TTCGTCTTTA TAGTCGAAGT TAACACCATT TACATTCTTC AGTCCTCTG CTACAGATGG	3480
CACTGTTGTC ACTACTGCCA CTTTATTATT TTTAGCCATA GAAGAACCTT GGAGATGCC	3540
AATTCCCTACA GAGATTCCTA AAAAGAGGAA CGGCAGAAC ACCATAAAGA AGAAACTCCA	3600
TGACTCGACA TGTGAAAGAT AGGTTCCCTT GATTACAACC CACATATTTC TCATAACTTCC	3660
ACTCCTGATT CTAGTTAAA GATTTCATCG ATAGTTGGCG CTTGTTGGTC AAATGTTGCG	3720
ATATATTGAC CTTGAGTCAA GATTGAGAAG AGTTCCCTTC CAGCGCTCTC ATCCTCCAAA	3780
ATCAATTTC AACTGCCCTTG TTTGGTCAAG CTCACCTGTT TGACATGAGG AAGATTTCC	3840
AATTCTCCCT TGCTTCGTTA ACTTGAAACA AAGAGACGCG TTTTCCCGTA TTGATTGCGG	3900
ACATCCTGAA CTGGTCCGTG CAAGACCACA CGGCCATCTC GGATCATCAG AATATCGTCA	3960
CAAAGTTCCCT CAACATTGGT CATGACATGG TCAGAAAAGA TAATGGTTGT CCGCGCTCTT	4020

321

TTTCCTGAAA AATGACTTGT TTGAGCAATT CTGTATTAAC TGGGTCCAAT CCACTAAAAG	4080
GCTCATCCAA GATAATCAGG TCTGGTTCAT GAATCAGAGT AATAATGAGC TGAATCTTCT	4140
GCTGATTTCC TTTTGACAGA CTCTTGATTT TATCTGTCAG CTTTCCTTTC ACTTCCAACC	4200
TCTTCATCCA TTGAGGGAGT TTTTCTTGA CTTCTTTGGC ATCCATGCCT TTTAGAGTCG	4260
CCAAGTAGCG AACTTGTCA AGAACTGTCA ATTTAGGCAT GAGATGCGTT CTTCAGGCAG	4320
ATAACCAATC CGAGCATAGG TCTCCTGACG AATATCCTGA CCATCCAGAC CGATTTCTCC	4380
CTGATATTCT AGGAATTTCAGA AATAACTATG GAAAATCGTT GTTTTCCAG CACCATTTT	4440
TCCGACTAGT CCCAAATAC GACCTGGTCG CGCTTGAAAG TCAATACCAA ACAAAACTTG	4500
CTTGGATCCA AAACCTTTCT CTAGACTTCT TACTTCTAGC ATCTTCACC TCCGAAATT	4560
CTTGCACCTCA TTATACTCCT TTTTGATAGC CTTTACAATG TTTTTGTCC ATTTTTAGAA	4620
GACTATPGCT GTGTAAGATA TGGCCTGGAG CACTTTTATA CTCAATGAAA ATCAAAGAGC	4680
AAACTAGGAA GCTAGCCGTA GACTGCTAA ACTACAGCTT TGAGGTTGCA GATAAAACTG	4740
ACGAAGTCgA CTCAAAACAC TGTGTTGAGG TTGTGGATAG AACTGACGAA kCrTAaCTAT	4800
ATCTACGGCA AGGCGAactG ACCTGGTTG AAGAGATTT CGAAGAGTAT TAGTGATAAA	4860
TCCATTATAC AGCAGCAAAC TTAATTATA CCTTCCGCTC CTCAACTGTC TATTTTTAAT	4920
CCTGAATTGT TATTGAGTA ACTCCCTTTT CCTCGTAAAG TTTTCTTCCT CTAAAACCTC	4980
TGGAAAAGG CTAATAGTTT CAGACAACTT TTTTATAAGA AACAAGTTCA TGTGTCATT	5040
CAAGAAGGAG TAATCCTTTA TCTACTAATG GACGGAACAG AATTCAACCG CTTGTCCGAT	5100
ATGTTTCTA AGGATTATAT AGTAAAATGA ATAAGAACCA GGACAAATG ATCAGGACAG	5160
TCAAATTGAT TTCTAAACAAT GTTTAGAAG TAGATGTATA CTATTCTAGT TTCAATCTGC	5220
TATATCTATT ATGCACACCC CTATAGGATC TAATGAAAAT CACAACAGGC TCATTCTAG	5280
ATGGTTACCT AAGCCTAAGG GAACTAAGAA AACGACTACC AAGGAAGTCG CATTCTCGA	5340
AAAGTAGATT AACAACTATC CTAAAAAATG CTTGAACACTAC AAGTCCCCCA GAGAAGACTT	5400
CTGGATGACT AACTGAACT TGAAATTAG CAATAATTAA TTCACTATCT AACTATATT	5460
AGTAATTATT TCAGAACTGA TTAATATTAA ATTAACCTAA CAATTCAAAG GATTCTACT	5520
AGCCATAAAAT TACGTCCATC AGAGAGAGAC TCTTACTACT TTTAGATTT AGTCTTTCTA	5580
GCTTCAGAAAT ACATCTAAAC TTTAGGGAAA ATGACTATTC GAAAGCGCGA ATGCCTAAA	5640
ATTATCTCAG ATAAGCTATT CGAAACTTAG AATGCTTTA AATTATGGA ATTGCGATTA	5700
TTCGAAACCT AGAATGCATA TAACCTTTAG TTGACAGACC TATTCTAAGT CTCGAAGGGC	5760

322

TATTTACTTT	CTATTCCTTA	TCAAAAAAGA	CTCATTCCCC	CTTTCTCCTC	CAAAATATGG	5820
TATAGTAGAA	ATATACTATC	TATGAGGAGT	TTACATGTCA	CAGGATAAAC	AAATGAAAGC	5880
TGTTTCTCCC	CTTCTGCAGC	GAGTTATCAA	TATCTCATCG	ATTGTCGGTG	GGGTTGGGAG	5940
TTTGTATTTTC	TGTATTTGGG	CTTATCAGGC	TGGGATTTA	CAATCCAAGG	AAACCCCTCTC	6000
TGCCTTTATC	CAGCAGGCAG	GCATCTGGGG	TCCACCTCTC	TTTATCTTT	TACAGATTTT	6060
ACAGACTGTC	GTCCCTATCA	TTCCAGGGGC	CTTGACCTCG	GTGGCTGGGG	TCTTTATCTA	6120
CGGGCACATC	ATCGGGACTA	TCTACAACTA	TATCGGCATC	GTGATTGGCT	GTGCCATTAT	6180
CTTTTATCTA	GTGCGCCTAT	ACGGAGCTGC	CTTTGTCCAG	TCTGTCGTCA	GCAAGCGCAC	6240
CTACGACAAG	TACATCGACT	GGCTAGATAA	GGGCAATCGT	TTTGACCGCT	TCTTTATTTT	6300
TATGATGATT	TGGCCCATTA	GCCCCAGCTGA	CTTTCTCTGT	ATGCTGGCTG	CCCTGACCAA	6360
GATGAGCTTC	AAGCGCTACA	TGACCATCAT	CATTCTGACC	AAACCCTTTA	CCCTCGTGGT	6420
TTATACCTAC	GGTCTGACCT	ATATTATTGA	CTTTTCTGG	CAAATGCTTT	GACACGTAAA	6480
AAATCCGTTT	GGTTTCCCAA	GTGGATTTTT	AAAGCGTAGA	TTAACTATAG	CTTGATACTA	6540
AATATACTTT	GGTATGGAAA	TCATGCATAT	TTTCGATAG	TGAGGCGAGG	ACTTACCTAG	6600
CCTTTCCGCC	GTGATAGAAA	CACCTGAAAT	CTAATGGTTT	CAGGTATTG	GAAACTTGA	6660
GCCTAGTGTG	TCAAAGTTA	GGTATGGAAT	TTTGAAGAAA	GTCGCTACCG	TCCGTAATCA	6720
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GATTTTGTGC	TTTATTTTGA	AACTTCTTTT	GCAAGAACAA	AGTTCCAAG	TGTGGCAGAA	6840
CCATTTCCTG	CGACTGCTGG	CGTCACGATA	TAGTCACGCA	CATCTGGTAC	TGGTAGGTAA	6900
CCATTAAGAA	GAGATGTAAA	TTTCTCACGG	ACACGGTCCA	GCATATGTTG	TTGAGCCATG	6960
ACCCCTCCAC	CAAAGACAAT	CACGTCTGGG	CGGAAAGTCA	CTGTCGCATT	AACCGCAGCT	7020
TGAGCGATAT	AGTAGGCTTG	AACATCCCAA	ACAGGGTTGT	TGAGTTCAAT	AGTTTCCCA	7080
CGTACACCTG	TACGAGCTTC	CAAACTTGGA	CCAGCTGCAT	AACCTCTAG	ACATCCCTTA	7140
TGGAAAGGAC	AAACACCCCTT	AAACTCTTTT	TCAATATCCA	TTGGGTGTCT	AGCAACATAA	7200
TAATGACCCA	TTTCAGGGTG	ACCCACACCA	CCGATAAACT	CACCACGTTG	GATGACGCC	7260
GCACCGATAC	CTGTACCGAT	TGTGTAGTAA	ACCAAGTTTT	CGATACGACC	ACCAGCATG	7320
TTACGGGCAA	CCATTTCACCC	GTAAAGCAGAG	CTGTTTACGT	CTGTTGTGAA	GTACATTGGC	7380
ACGTTTAGGG	CGCGACGAAG	GGCACCAAGC	AAGTCTACAT	TTGCCAGTT	TGGTTTTGGA	7440
GTCGTCGTGA	TAAAGCCATA	AGTTTTGAG	TTTTGTCAA	TATCAATCGG	CCCAAATGAA	7500
CCAACTGCAA	GACCAGCAAG	GTTATCGAAT	TTTGAGAAGA	ACTCAATGGT	TTTATCGATT	7560

323

GTTCGATTG GAGTTGTTGT TGGAAATTGT GTTTTTCTA CAACGTTAAA GTTTTCATCA	7620
CCGACAGCAC AGACAAACTT TGTACCGCCC GCTTCCAAGC TTCCATATAA TTTTGTCTAG	7680
ATAAACCTCT TGTGTTTATT TTCTTTATTA TAGCATACTT CGAAAGTCTA AATGTCTCTA	7740
TTTTTAGAT TTCCCTCTGT AAATCTTACT ATCTAATAAA AACGAACAAA CATGTCATT	7800
GTTCGTTTC ACATTAGAGA GGATTGATTA GATTTTCACT TCGATCACAG CATCCCCCTT	7860
AGCAACTGAA CCTGTTGCGA CTGGAGCTAC TGAAGCGTAG TCACCTGTAT TTGTAACGAT	7920
AACCATTGTT GTATCATCAA GTCCAGCTGC AGCGATTTG TTTGAGTCAA ATGTTCCAAG	7980
AACATGCCA GCTTCACCT TATTACCTTG AGCAACTTTT GTTTCAAAAC CGTCACCGTT	8040
CATAGATACA GTATCAATAC CAACATGAAT CAAACTTCA GCACCATTTC TTGTTTCAA	8100
ACCAAAAGCG TGCCCTGTTG GAAAGGCAAT TGAAACTTCA GCATCAGCTG GTGCATAGAC	8160
CACGCCCTGG CTTGGTTCA CAACGATACC TTGTCCTCATA GCTCCACTTG AGAAGACTGG	8220
GTCATTGACA TCAGCAAGAG CGACAAACATC ACCGACGATA GGAGTTACAA GTGTTTCATT	8280
TTGAAGAGCT GCTGGCAGA CTTCTTCTTT TTCTTCAGCC ACTTCAGCTC GTTTTGCAGC	8340
TGCAGTTGCG TCTACTTCAT CTTCGTAACC AAACATGTAA GTAAGAGCAA AACCAAGGGC	8400
AAATGATACA GCTACCATAA GAAGGTATTG TGGAAGTTGT CCGTTACCAA CATAAAGCAT	8460
TGTACCAGGG ATGATGGTGA TACCAATTACG AGTACCGACA AGTCCAAGGA TAGAAGCCAA	8520
TCCACCACCG ATTGCACCG CAATCAATGA AAGGAAGAAT GGTTCACGGA AGCGCAAGTT	8580
CACCCCGAAG ATAGCAGGCT CTGTAATACC TAGGAAGGCA GAAAGAGCAG CCGGGAAAGC	8640
AAGTGTGTTTC AGTTTGTGAT TTTTGTGTTT AACACCAACC GCAACAGTAG CAGCACCTTG	8700
AGCTGTCATA GCAGCTGTGA TGATAGCGTT GAATGGGTTA GCATGGTCAG CAGCAAGTAA	8760
TTGCACTTCA AGCAAGTGA AGATGTGGTG CACACCTGAC ACGACGATCA ATTGGTGAAC	8820
CCCACCAATC AAGAAACCAC CAAGACCAAA TGGCATGCTA AGAACATCGTT TTGTAGCAAT	8880
AAGGATGTAG TTTCAACAA CGTGGAAACAC TGGTCCAATG ACAAAAGAGTC CAAGGATAGA	8940
CATGACCAAA AGTGTACGA ATGGTGTAC CAAGAGGTCA ATGACATCTG GAACAACTTG	9000
CGGACAGCTT TTTCAAATT AGCTCCGACA ACCCCGATGA TGAAGGCTGG AAGAACGGAA	9060
CCTTGCAACAC CAACACAGG GATGAAACCA AAGAAGTTCA TCGCTGTTAC TTCACCACCT	9120
TGAGCAACTG CCCAAGCGTT TGGAAGTGAG CCAGAGACAA GCATCATAACC AAGAACGATA	9180
CCAACGGCAG GATTCCACC AAATACACGG AAGGTTGACC ACACAACCAA ACCTGGCAAG	9240
ATGATGAAGG CTGTATCTGT CAAGATTGTG GTGTAAGTTG CAAAGTCACC TGGAAGTGGC	9300

324	
ATTTCAAGAG CGTTGAAAAG ACCACGCACA CCCATGAAGA GACCTGTCGC TACGATAACT	9360
GGGATGATTG GAACGAAAAC ATCACCAAAA GTACGGATAG CACGTTGGAA CCAGTTCCCT	9420
TGTTTAGCAA CTTCTGCTTT CATGTCATCC TTAGATGATG TTGGTAATCC AAGTACAACA	9480
ACTTCATCGT ACATTTGTT AACTGTACCT GTACCAAAGA TAATTTGGTA TTGCCCTGAG	9540
TTAAAGAAAG CACCTTGAAC TTTTCAGAAG TTCTCAATCA CTTCTTTATT GATTTCTCT	9600
TCATCTTGA CCATGACACG TAGACGAGTC GCACAGTGGG CAACACTATT GACATTTCA	9660
CGTCCGCCCA AGGCATCGAT GACTTTTTTT GCAATTTCCT GATTGTTCAT TTGCAAAAT	9720
CTCCTTATAT AACATTTGTT TCTTGTGTTGA AAGCGATTTT ATTGCGCCG	9769

(2) INFORMATION FOR SEQ ID NO: 31:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 3149 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 31:

CGCTTGAGTG CTAATTCTATA GTTCTATTGT ATCACTTGGT CAGAAATAAT CAAGAAAAAA	60
GTCTGACTTT CTCAAGATAA AAAGCCTGAG ACCAACTCG ACTTTTTAAT TCTTAAAATG	120
GCAATTCTTC CTCTTCCAAG ACCAAATCTG CCAAATCTG GCCTGCATTA TTTTCACGCA	180
TAGCACGTTT GGCACGACTT TCCAAGAGTT GGAATCCTGT GACAAGTACT TCGGTACAGT	240
AGTTCATTTG GCCATTTTC TCAAAGCGAC GGGTACGAA TTCTCCATCA ACGGAAATGA	300
GACTACCTTT GGTTGCGTAC TTGCCAAAGT TTCTGCTAGT CTGCCCTATA GGACCATATT	360
GACAAATCA GCTTCACGTT CACCGTTTG GTCTTGTAA CGACGGTTCA CAGCGATAGT	420
TGCTCGCGCT ACCGACTTGT CATTGTTGGT TTTGTGCAAT TCTGGTGTAG ACGTTAAACG	480
TCCAATCAAG ATAACCTTAT TATACATATT TTCTCCCTCC TACTTATCTA TTCTGAGGAA	540
ATCAAAAAAA GTTACAGAAA TTGTAACTT TTGAGAAAA TTTTTTATTT TTTATGAACC	600
ATGAAACCTG TCGCCTGTTG ATTGGCCATA ATGGTCATAT CTGTAATCTG AACACGACGA	660
GGTTGACTAG TCACATAGAC TACTGTATCT GCAATATCCT GAGCTGCAA AGCTTCTATT	720
CCTTGGTAA CGGACGCAGC TCGTTCTTA TCACCATGAA AACGCAGTGT AGAAAAATCT	780
GTTCGACAA TTCCAGGCTG AATGGTCGTC ACCTTGATAT CCGTTGCGAT GGTATCAATT	840
CGCAGTCCAT CTGAAAAGGT CTTAACTGCC GCCTTGGTGG CTGAGTAAAC AGCTGCACCA	900
GCATAGGCAT AAATTCCCTGC GGTTGACCCC ATATTGATAA TATGACCTTG ATTGGCTTT	960

325

ACCATTGCTG GCAAGAAACA GCGAGTGACT GCCATCAAAC	CTTGACATT GGTATCCAAC	1020
ATGGTCAGCA TATCCAACTC TTCATAGTCT TGATAGGGAG	CTAAGCCAAG AGCCAGTCCT	1080
GCGTTATTGA CCAGGATGTC AATCTGACCT ATCGTTCTA	AAATATCAGA GCAGACAGTC	1140
TTTACCATTG TCATATCCGT GACATCTAGG AGAAAAGTCC	AAACTGTTG ATTTGGAAA	1200
GTTTCTGCAA ACTCCGCCTT AAGAGCTCT AGTCTGTCTA	TCCGTCGTC TGTAGAACG	1260
ACATCCTCAC CCTGCTCCAG ATAAGCACGC GCAATCGCTT	CACCGATTCC TGATGTCGCT	1320
CCTGTAATCA CAACATTTT TGCCATCTA TTTCCTCTA	GCTGGTCTAT CAGATATTAA	1380
CAACTCTTA GGCAGTCCAG TCTTCGCTG GTCGACGG	TGTTCCGACA ACTTGGTCTT	1440
CTGATAATTC AAGCACCCCA CGTTTTGCTG GAGCATTGG	CAGATGCAAT TCACGAGGAC	1500
TGCACATCAT ACCAAAAACTC TTTTCAACCAC	GAAGTTCACC TGGGAAAATG	1560
TTGGCATCAT AGCTCCAGGA AGCGCGACAA TGTTTTCAA	CCCCACACGC GCATTGGAG	1620
CTCCTGCAAC GATTTGTACA CTCTTATCAC TTGCGACTGC	AACTTGGCAG ATGTTGAGGT	1680
GGTCACTATC TGGATGGGCT ACCATCTCAA CAATTCACC	TACAACAAAC TTAGGTTCT	1740
TATCATTAAC AATTTCTCT GTAAAACCTT CCGCCTGCAA	CTCTTGGTTC AAACGAGCGA	1800
CTTGCTCATC TGTCAAAAG ACTTGACCGC GCTCTGCAAT	TTCAAATAAA CTTGAAACTT	1860
CGAAAATATT CCAAGCCACT GTTCCCCAT TATCTTGAG	AAAAACACGG GCTACCTTGC	1920
CTTTGGCCTC CACATCCACT TTGGCATCTC CGCTATTTT	CACGATGACCA ATAAGGACAT	1980
CACCGACATG TTCTTATTA TATGTAAAAA TCATTGTTTC	CTTTTCTCC TATTTCAGTC	2040
CTGCTAAAAA GTCATTGATT TTGTTGCTTGC TTTTACGGTC	GCGATTGACA AAACGACCGA	2100
TTTCCTGTC CTTTCTAGA ACAACAAGGC TAGGAATTCC	GTAAACATCC CAGAGTTGG	2160
CCAATCCAT ATACTGATCT CGGTCCATTG	GAATAAGGT GAACTCTGGA TTGGTCTCCT	2220
CAATCTCTGG TAAGGCAGGA TAAATATAAC GACAATCGCT	ACACCACTCT GCCACAAAAA	2280
TGAAGACCTT CTTGCCGCT TTTCCACTA AAGATGCTAA	TTCTTCTAAA CTTGCTGGCT	2340
GTATCATAAG ACTTCCCTC CATAGACTAG GTCTTCATTT	TCATAGACAA AGGTATAATG	2400
ACGGCCATCC TCAAAATGA CGCCACCAAC CAAGCTCTCC	AGACTGCTTT CGTAAACTTG	2460
AACATAAAGG GTCGCAATTG CCCCCATGTC GGAAAAATGG	TCTCGCACAA TCTCTGTCAA	2520
CTCTTCTCTGA GTCTTCATGA GCTTACGGTC ATCTGCAACT	TTTTTCTGAG CAAGAGCAAAG	2580
GCTTCCGATA CCTAGCAGAG CCAAGCCTGC CATCCACATT	TTTTTAGCTT TCATACCATT	2640
CATTTTAACA CAAAAAAGGC TTCAGGACAA ATGAGGAAGC	AGCAGAAAAG CAAGTAAAAA	2700

326	
GCCTCTTCCCT TTAAGGAAAA GGACTTCTTA TACTCAATGA AAATCAAAGA CCAAACCTAGG	2760
AAGCTAGCCG CAGGCTGCTC AAAGCACTGC TTTGAGGTTG TAGATAGAAC TGACGAGTCa	2820
CTCAAAACAC TGGTTTGAGG TTGAGGATGA AGCTGACGTG GTTGAAGAG ATTTTCGAAG	2880
AGTATTATTC TTATTGCCAG GCACCTAAGT TGCCAACGTA GTAACATATCA GGTGTGTAGG	2940
TATTGCGAGC ATCTTACCTG ATGAAGCCAG ATAATAC TGTCCATTGT CTTTGACCCA	3000
ATCATTCGCA ATCATGGAAC CAGAAGAACT TACATAATAC CATTCTCCCT TGTCTAAAC	3060
CCAAGTACTG ACTTCATGG TTCCGTGAGCA ATTAAAGGCA AAAAAACTGT CCAATAACAT	3120
TCGTTTTTA AAAGCATTG ACACATAC	3149

(2) INFORMATION FOR SEQ ID NO: 32:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 10240 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 32:

CCAAAAAATTC AACCTTTAAG GGGAGTCCAG AGAGACTCAC AAGGTGTCAG ATAAAAGAAT	60
GGTGCAATT TCTAGAGGAG ACTTTTGAG TGTGCTCTCT TGTGTTGTAC GATTTTAAC	120
GAGGCCTTGC ACTAGCAAGG TCTTTCTTT ATCTGGTCCC CTTAAAATTT AAGGAGGAAA	180
AGTTATGAAT CCCACATGTA AGAAGCGTTT GGGTGTCAATT CGGTTGGAAA CCATGAAGGT	240
GGTTGCACAA GAGGAAATCG CGCCACAATC TTTGAATTAG TCCTAGAAGG AGAAATGGTT	300
GAAGCCATGC GAGCAGGCCA ATTCTTCAT CTGCGTGTAC CGGACGATGC CCATCTTTA	360
CGTCGTCTTA TTTCAATTTC GTCTATTGAC AAGGCAAACA AGCAGTGTCA CCTCATTAT	420
CGGATTGACG GAGCTGGGAC TGCAATTTC TCAACCTTAA GTCAGGGAGA CACTCTTGAT	480
GTGATGGGGC CTCAGGGAAA TGGTTTGAC TTGCTGACC TTGATGAGCA GAATCAGGTT	540
CTCCTTGTG TGTTGGGAT TGGTGTCCA CCCTGCTTG AGGTGGCCAA GGAATTGCAT	600
GAACGTGGAG TGAAAGTAGT GACAGTCCTC GTTGTGCTA ATAAGGATGC TGTTATTG	660
AAAACGGAAT TGGCTCAGTA TGGTCAGGTC TTTGTAACGA CAGATGATGG TTCTTATGGC	720
ATCAAGGGAA ATCTTCCGT TGTATCAAT GATTAGACA GTCAGTTGA TGCTGTGTTAC	780
TCGTGTGGGG CTCCAGGAAT GATGAAGTAT ATCAATCAA CCTTGATGTA TCACCCAAGA	840
GCCTATTTAT CTCTGGAATC TCGTATGGCT TGTGGGATGG GAGCTTGCTA TGCTGTGTT	900
CTAAAAGTAC CAGAAAACGA GACGGTCAGC CAACGCGTCT GTGAAGATGG TCCTGTTTC	960

CGCACAGGAA CAGTTGTATT ATAAGGAGAA AATTATGACT ACAAATCGAT TACAAGTTTC	1020
TCTACCTGGT TTGGATTGAA AAAATCCGAT TATTCCAGCA TCAGGCTGTT TTGGCTTGG	1080
ACAAGAGTAT GCCAAGTACT ATGATTTAGA CCTTTTAGGT TCTATTATGA TCAAGGCAGAC	1140
AACCCCTGAA CCACGTTTG GGAATCCAAC TCCAAGAGTG GCAGAGACGC CTGCTGGTAT	1200
GCTCAATGCA ATTGGCTTGC AAAATCCTGG TTTAGAGGTT GTTTGGCTG AAAAGCTACC	1260
TTGGCTGGAA AGAGAATATC CAAATCTTCC TATTATTGCC AATGTAGCTG GTTTTCAAA	1320
ACAAGAGTAT GCAGCTGTT CTCATGGGAT TTCCAAGGCA ACTAATGTAA AAGCTATCGA	1380
GCTCAATATT TCTGTCCCCA ATGTTGACCA CTGTAATCAT GGACTTTGA TTGGTCAAGA	1440
TCCAGATTTG GCTTATGATG TGGTGAAGC AGCTGTGGAA GCCTCAGAAG TGCCAGTTA	1500
TGTCAAATTA ACCCGAGTG TGACCGATAT CGTTACTGTC GCAAAAGCTG CAGAAGATGC	1560
GGGAGCAAGT GGCTTGACCA TGATCAATAC TCTGGTTGGA ATGCGCTTTC ACCTCAAAAC	1620
TAGAAAACCA ATCTTGGCCA ATGGAACAGG TGGAAATGTCT GGTCCAGCAG TCTTCCAGT	1680
AGCCCTCAAA CTCATCCGCC AAGTTGCCA ACAACAGAC CTGCCATATCA TTGGAATGGG	1740
AGGAGTGGAT TCGGCTGAAG CTGCCCTAGA AATGTATCTG GCTGGGGCAT CTGCTATCGG	1800
AGTTGGAACA GCTAACTTTA CCAATCCTTA TGCCTGCCCT GACATCATCG AAAATTAC	1860
AAAAGTCATG GATAAAATACG GTATTAGCAG TCTGGAAGAA CTCCGTCAGG AAGTAAAAGA	1920
GTCTCTGAGC TAAACTGCAA TCAATCTGTT CTTGATTTTT TATTAGTTTG TAATATGAAT	1980
TTAGGAGAAT TTTGGTACAA TAAAATAAT AAGAACAGAG GAAGAAGGTT AATGAAGAAA	2040
GTAAGATTTA TTTTTTTAGC TCTGCTATTT TTCTTAGCTA GTCCAGAGGG TGCAATGGCT	2100
AGTGATGGTA CTTGGCAAGG AAAACAGTAT CTGAAAGAAG ATGGCAGTCA AGCAGCAAAT	2160
GACTGGGTTT TTGATACTCA TTATCAATCT TGGTTCTATA TAAAAGCAGA TGCTAACTAT	2220
GCTGAAAATG AATGGCTAAA GCAAGGTGAC GACTATTTT ACCTCAAATC TGGTGGCTAT	2280
ATGGCCAAAT CAGAATGGGT AGAACAGAAC GGAGCCTTTT ATTATCTTGA CCAAGATGGA	2340
AAGATGAAAA GAAATGCTTG GGTAGGAAC TCCATGTTG GTGCAACAGG TGCCAAAGTA	2400
ATAGAAGACT GGGTCTATGA TTCTCAATAC GATGCTTGTT TTTATATCAA AGCAGATGGA	2460
CAGCACGCAG AGAAAGAATG GCTCCAAATT AAAGGGAAGG ACTATTATTT CAAATCCGGT	2520
GGTTATCTAC TGACAAGTCA GTGGATTAAT CAAGCTTATG TGAATGCTAG TGGTGGCAA	2580
GTACAGCAAG GTTGGCTTT TGACAAACAA TACCAATCTT GTTTTACAT CAAAGAAAAT	2640
CGAAACTATG CTGATAAAAGA ATGGATTTTC GAGAATGGTC ACTATTATTA TCTAAAATCC	2700

328	
GGTGGyTACA TGGCAGCCAA TGAATGGATT TGGGATAAGG AATCTTGGTT TTATCTCAAA	2760
TyTGATGGGA AAATrGCTGA AAAAGAATGG GTCTACGATT CTCATAGTCA AGCTTGGTAC	2820
TACTTCAAAAT CCGGTGGTTA CATGACAGCC AATGAATGGA TTTGGGATAA GGAATCTTGG	2880
TTTTACCTCA AATCTGATGG GAAAATAGCT GAAAAAGAAT GGGTCTACGA TTCTCATAGT	2940
CAAGCTTGGT ACTACTTCAA ATCTGGTGGC TACATGGCGA AAAATGAGAC AGTAGATGGT	3000
TATCAGCTTG GAAGCGATGG TAAATGGCTT GGAGGAAAAA CTACAAATGA AAATGCTGCT	3060
TACTATCAAG TAGTGCCTGT TACAGCCAAT GTTTATGATT CAGATGGTGA AAAGCTTCC	3120
TATATATCGC AAGGTAGTGT CGTATGGCTA GATAAGGATA GAAAAAGTGA TGACAAGCGC	3180
TTGGCTATTA CTATTCTGG TTTGTCAGGC TATATGAAA CAGAAGATTT ACAAGCGCTA	3240
GATGCTAGTA AGGACTTTAT CCCTTATTAT GAGAGTGTAGT GCCACCGTTT TTATCACTAT	3300
GTGGCTCAGA ATGCTAGTAT CCCAGTAGCT TCTCATCTTT CTGATATGGA AGTAGGCAAG	3360
AAATATTATT CGGCAGATGG CCTGCATTTT GATGGTTTTA AGCTTGAGAA TCCCTTCCTT	3420
TTCAAAGATT TAACAGAGGC TACAAACTAC AGTGCTGAAG AATTGGATAA GGTATTTAGT	3480
TTGCTAAACA TTAACAATAG CCTTTTGGAG ACAAGGGCG CTACCTTTAA GGAAGCCGAA	3540
GAACATTACC ATATCAATGC TCTTTATCTC CTTGCCATA GTGCCCTAGA AAGTAACTGG	3600
GGAAGAAGTA AAATTGCCA AGATAAGAAT AATTCTTTG GCATTACAGC CTATGATACG	3660
ACCCCTTACC TTTCTGCTAA GACATTTGAT GATGTGGATA AGGGAATTAA AGGTGCAACC	3720
AAGTGGATTA AGGAAAATTA TATCGATAGG GGAAGAACTT TCCTTGGAAA CAAGGCTTCT	3780
GGTATGAATG TGGAAATATGC TTCAGACCTT TATTGGGGCG AAAAAATTGC TAGTGTGATG	3840
ATGAAAATCA ATGAGAAGCT AGGTGGCAA GATTAGTACT ATAAGTGAAT ATGATTTGAG	3900
TGAATAGTAA GTTAAAATC CTGATTTCAA GTAAAATCAG GATTTTTCA TGGATGCAAT	3960
TTTTTTGGAG TCTGGTGTGA CGCGGAGGGT CTTTTGTCT GTGTAAGTGA CAAAGCCGGG	4020
TTTTCCACCA GTGGTTTAT TGAGTTTTT GACTTCATTC ATATCTACCT GCACCAGATT	4080
CGACAGGCAGC CCTTGAGAGA AGTAGGCAGC TAACTCTGCT GCGTCTGTCT TGACTGCATC	4140
AGATGGGTCA AGATTCCTG AGATGACAAC ATGGCTTCCA GGAATGTCT TAGCATGGAA	4200
CCAAAGTTCC TCCTTGCGGG CCATTTAAA GGTCAATTCC TCATTTGAA GATTGTTTCG	4260
TCCGACATAG ATGATGGTTT TGCCATCGCT TGCTAGATAT TGTTCTAGTT TTTTGGCTTT	4320
CTGGATTTTC TCCCCTGTC TTCTGCGGAT AAAACCTGTT TGAATCAATT CTTCACGGAT	4380
TTCAGCGATT TCTTCCAGTC CAGCTTGGTT GAGGACGGTT TCTACACTTT CCAGATAGAG	4440
AATAGTGGCT TTGGTTCTT CAATCAAATC AGTCAAGTAT TTGACAGCTT CTTTGAGTTT	4500

329

CTGATACCGT TTAAAATAGC GTTGGGCATT CTGGTTGGGA GTCAGAGCCT TATCAAGCGC	4560
AATCATGATA GGTTGGTTGG TATAGTAGTT GTCTAGGATA ACCTGGTCTT GGTCGTTAGG	4620
CACTTGGTGG AGGAAGGTTG TCAGCAATT TCCTTTTGA CGAAATTCTT CAGCGTTGTC	4680
TGTGCCAGT AACTTTTTT CCTGTTTTT GAGTTGTGT CGGTTTTCTT GAAGTTTATT	4740
TTCAACACGA CGAACAGTT CACTGGCTG CTGTTGACG CGGTCGCGCT CAGCCTTATC	4800
CTTATAGTAG GTGCCAACA AATCAGAAAG ATTTGCAAAA GGCTCTCCCA CCTGATTG	4860
AAAAGGAACG GGACTGAAGG AAGTCTCAGT CAAGCATGGC TTGGTTTCTT GATTGAAAAA	4920
ATTTGGAAA GCGGAAAGTT TTTCACTAAC CAGTATCCTT TCCAATTCTAT TTGCGTATC	4980
GCGTCCCAGA CCTTGAAGA GGCTTGAAAG ATTTTTGCT GTTAGTTCTT GGGTTGCAG	5040
GATTCAAAG AGCTTTTCACT CCTTGATAGT AAAAGGATTG AGAGATTTG TACTGGCGG	5100
AGCGATATAG GTCGATCCTG GAAGTAAAGT GCGGTAGCTA TTTTGTGAAA AGCCGACGTG	5160
TTTGATAACT TCGAGGATT TATGACTGCT TTTATCGACC AGTAGAATAT TACTGTGTT	5220
CCCCATAATT TCGATAATCA AGGTAGCCTG GATATGGTCT CCAATCTCGT TTTTATTGGA	5280
AACTGTAATT TCCACAATAC GGTCAATTTC CACTGCTCA ATCGACTCAA TCAGGGCCCC	5340
CTGCAAATAC TTTCTCAAAA CCATGATAAA GGTAGAAGGT TGAGCTGGAT TTTCAAAGT	5400
CGTTGGGTC AGCTGAATGC GTCCAAAAAC TGGATGGCA GAAAGGAGCA GGCGATGGCT	5460
TTGGCGATTG CTGCGGATTT GCAAGACCAA CTCTGGTCA AAAGGCTGAT TGATTTCTG	5520
GATGCCACCA TTCACTAATT CGCTTCGCAA TTCCCTCAACT ATGTGGTGA AAAAAAATCC	5580
GTCAAATGAC ATCGTTCTCT CCTTGATGATT GTATTCCATA GTATTATATC AAAAAAGGTAG	5640
AATAAAATCA TGGAAATGTG GTATAATAAA GCCAAGTAAA GAGAAACGAG AAGCACATGT	5700
ATATTGAAAT GGTAGATGAA ACTGGTCAAG TTTCAAAAGA AATGTTGCAA CAAACCCAAG	5760
AAATTTGGA ATTTGCAGCC CAAAATTAG GAAAAGAAGA CAAGGAGATG GCAGTCACCT	5820
TTGTGACCAA TGACCGTAGT CATGAACCTTA ATCTGGAGTA CCGTAACACC GACCGTCCGA	5880
CAGATGTCACT CAGCCTTGAG TATAAACCAG ATTTGAAAT TGCCCTTGAC GAAGAGGATT	5940
TGCTTGGAAA TTCAAGAATTG GCAGAGATGA TGTCTGAGTT TGATGGCTAT ATTGGGGAT	6000
TGTTCATCTC TATCGATAAG GCTCATGAGC AGGCCGAAGA ATATGGTCAC AGCTTGAGC	6060
GTGAGATGGG CTTCTTGGCA GTACACGGCT TTTTACATAT TAACGGCTAT GATCACTACA	6120
CTCCGGAAGA AGAACGGAG ATGTTGGTT TACAAGAAGA AATTTGACA GCCTATGGAC	6180
TCACAAGACA ATAAACGAAA ATGGAAAAAT CGTGACTTGA TATCCAGTTT AGAATTTGCT	6240

330

TTGACAGGTA TTTTACTGC TATCAAGGAA GAACGCAATA TGCAGAAACA CGCAGTGACG 6300
 GCTCTAGTGG TCATCCTTGC AGGTTTGTT TTTCAGGTGT CACGAATCGA ATGGCTCTT 6360
 CTCCTATTGA GTATTTCTT GGTAGTAGCC TTTGAGATTA TCAACTCTGC TATTGAAAAT 6420
 GTGGTGGATT TGGCCAGTCA CTATCACTTT TCCATGCTGG CTAAAATGC CAAGGATATG 6480
 CGGGCCGGCG CGGTATTAGT GGTTCTCTT TTCGCAGCCT TAACAGGCGC ATTGATTTT 6540
 CTCCCACGAA TCTGGGATT ATTATTTAA ACAGTAAGAG GAAATTATGA CTTTAAATC 6600
 AGGCTTTGTA GCCATTTAG GACGTCCAA TGTGGGAAG TCAACCTTT TAAATCACGT 6660
 TATGGGCAA AAGATTGCCA TCATGAGTGA CAAGGCGCAG ACAACGCGCA ATAAAATCAT 6720
 GGGAAATTAC ACGACTGATA AGGAGCAAAT TGTCTTATC GACACACCAG GGATTCAAA 6780
 GCCTAAACAA GCTCTCGAG ATTTCATGGT TGAGTCTGCC TACAGTACCC TPCGCGAAGT 6840
 GGACACTGTT CTTTCATGG TGCTGCTGA TGAAGGGCGT GGTAAGGGGG ACGATATGAT 6900
 TATCGAGCGT CTCAAGGCTG CCAAGGTTCC TGTGATTTG GTGGTGAATA AAATCGATAA 6960
 GGTCCATCCA GACCAGCTCT TGTCAGAT TGATGACTTC CGTAATCAA TGGACTTTAA 7020
 GGAAATTGTT CCAATCTAG CCCTTCAGGG AAATAACGTG TCTCGTCTAG TGGATATTTT 7080
 GAGTGAAAAT CTGGATGAAG GTTCCAATA TTTCCCGTCT GATCAAATCA CAGACCATCC 7140
 AGAACCTTTC TTGGTTTCAG AAATGGTTCG CGAGAAAGTC TTGCACCTAA CTCGTGAAGA 7200
 GATTCCGCAT TCTGTAGCAG TAGTTGTTGA CTCTATGAAA CGAGACGAAG AGACAGACAA 7260
 CGTTCACATC CGTGAACCA TCATGGTCGA GCGCGATAGC CAAAAAGGGGA TTATCATCGG 7320
 TAAAGGTGGC GCTATGCTTA AGAAAATCGG TAGCATGCC CGTCGTGATA TCGAACTCAT 7380
 GCTAGGAGAC AAGGTCTTCC TAGAAACCTG GGTCAAGGTC AAGAAAAACT GGCGCGATAA 7440
 AAAGCTAGAT TTGGCTGACT TTGGCTATAA TGAAAGAGAA TACTAAGTAG AGGTAGGCTC 7500
 ATGCCCTGCTT CTTGTTTTA CAGAAGGAGG ACTTATGCCT GAATTACCTG AGGTTGAAAC 7560
 CGTTTGTGCT GTGGTAAAGA AATTGATTAT AGGAAAGAAG ATTTCGAGTA TAGAAATTG 7620
 CTACCCCAAG ATGATTAAGA CGGATTTGGA AGAGTTCAA AGGGATTGC CTAGTCAGAT 7680
 TATCGAGTCA ATGGGACGTC GTGGAAAATA TTTGCTTTT TATCTGACAG ACAAGGTCTT 7740
 GATTTCCCAT TTGCGGATGG AGGGCAAGTA TTTTACTAT CCAGACCAAG GACCTGAACG 7800
 CAAGCATGCC CATGTTTCT TTCATTTGA AGATGGTGGC ACGCTTGTGTT ATGAGGATGT 7860
 TCGCAAGTTT GGAACCATGG AACTCTGGT GCCTGACCTT TTAGACGTCT ACTTTATTTC 7920
 TAAAAAATTA GGTCTGAAC CAAGCGAACCA AGACTTTGAT TTACAGGTCT TTCAATCTGC 7980
 CCTTGCCAAG TCCAAAAGC CTATCAAATC CCATCTCCTA GACCAGACCT TGGTAGCTGG 8040

ACTTGGCAAT ATCTATGTGG ATGAGGTTCT CTGGCGAGCT CAGGTTCATC CAGCTAGACC	8100
TTCCCAGACT TTGACAGCGAG AAGAAGCGAC TGCCATTCAAT GACCAGACCA TTGCTGTTT	8160
GGGCCAGGC GTTGAAAAAG GTGGCTCCAC CATTGGACT TATAACCAATG CCTTTGGGA	8220
AGATGGAAGC ATGCAGGACT TTCATCAGGT CTATGATAAG ACTGGTCAAG AATGTGTACG	8280
CTGTGGTACCT ATCATTGAGA AAATTCAACT AGGCGGACGT GGAACCCACT TTTGTCCAAA	8340
CTGTCAAAGG AGGGACTGAT GGGAAAATC ATCGGAATCA CTGGGGAAT TGCCCTGTT	8400
AAGTCAAATG TGACAAATTT TCTAACAGAC CAAGGCTTTC AAGTAGTGGG TGCCGACGCA	8460
GTCGTCCACC AACTACAGAA ACCTGGTGGT CGTCTGTTG AGGCTCTAGT ACAGCACTTT	8520
GGGCAAGAAA TCATTCTGA AAACGGAGAA CTCAATCGCC CTCTCCTAGC TAGTCTCATC	8580
TTTTCAAATC CTGATGAACG AGAATGGTCT AAGCAAATTC AAGGGGAGAT TATCCGTGAG	8640
GAACGGCTA CTTTGAGAGA ACAGTTGGCT CAGACAGAAAG AGATTTCTT CATGGATATT	8700
CCCCTACTTT TTGAGCAGGA CTACAGCGAT TGGTTGCTG AGACTTGGTT GGTCTATGTG	8760
GACCGAGATG CCCAAGTGGG ACCTGTTAATG AAAAGGGACC AGTTGTCCAA AGATGAAGCT	8820
GAGTCTCGTC TGGCAGCCC GTGGCCTTTA GAAAAAAAGA AAGATTTGGC CAGCCAGGTT	8880
CTTGATAATA ATGGCAATCA GAACCAGCTT CTTAACATCAAG TGCAATATCCT TCTTGAGGG	8940
GGTAGGCAAG ATGACAGAGA TTAACGGAA GGATAATCTG CGCATTGCCG GTTTGGTAA	9000
TTTTCTGACA GGAGCCAGTA TTTCTTGGT TGTACCTTTT ATGCCATCT TCGTGGAAA	9060
TCTAGGTGTA GGGAGTCAGC AAGTCGCTTT TTATGCAGGC TTAGCAATTCT GTCTCTG	9120
TATTTCCCGC GCGCTCTTTT CTCTTATTTG GGGTATTCTT GCTGACAAAT ACGGCCGAAA	9180
ACCCATGATG ATTGGGGCAG GTCTTGCTAT GACTATCACT ATGGGAGGCT TGGCCTTTG	9240
CCCAAATATC TATTGGTTAA TCTTCTTCG TTACTAAAC GGTGTATTG CAGGTTTTGT	9300
TCCTAACATCA ACGGCACTGA TAGCCAGTC GGTCCAAAG GAGAAATCAG GCTCTGCC	9360
AGGTACTTTG TCTACAGGCG TAGTTGCAGG TACTCTAACT GGTCCCTTTA TTGGTGGCTT	9420
TATCGCAGAA TTATTTGGCA TTGCTACAGT TTTCTTACTG GTTGGTAGTT TTCTATTTT	9480
AGCTGCTATT TTGACTATTT GCTTATCAA GGAAGATTTT CAACCAAGTAG CCAAGGAAA	9540
GGCTATTCCA ACAAAAGGAAT TATTACCTC GGTAAATAT CCCTATCTTT TGCTCAATCT	9600
CTTTTAACC AGTTTTGTCA TCCAATTTTC AGCTCAATCG ATTGGCCCTA TTTGGCTCT	9660
TTATGTACGC GACTTAGGGC AGACAGAGAA TCTTCTTTT GTCTCTGGTT TGATTGTGTC	9720
CAGTATGGGC TTTTCCAGCA TGATGAGTGC AGGAGTCATG GGCAAGCTAG GTGACAAGGT	9780

332	
GGGCAATCAT CGTCTCTTGG TTGTCGCCCA GTTTTATTCA GTCATCATCT ATCTCCTCTG	9840
TGCCAATGCC TCTAGCCCCC TTCAACTAGG ACTCTATCGT TTCCCTTTG GATTGGGAAC	9900
CGGTGCCTTG ATTCCCAGGG TTAATGCCCT ACTCAGCAAA ATGACTCCCA AAGCCGGCAT	9960
TTCGAGGGTC TTTCCTTCA ATCAGGTATT CTTTATCTG GGAGGTGTTG TTGGTCCCAT	10020
GGCAGGTTCT GCACTAGCAG GTCAATTGG CTACCATGCT GTCTTTATG CGACAAGCCT	10080
TTGTGTTGCC TTTAGTTGTC TCTTTAACCT GATTCAATTG CGAACATTAT TAAAAGTAAA	10140
GGAAATCTAG TGCGAGTAAA AATCAATCTC AAATGCTCCT CTTGTGGCAG TATCAATTAC	10200
CTAACCAAGTA AAAATTCAAA AACCCATCCA GACAGATTGA	10240

(2) INFORMATION FOR SEQ ID NO: 33:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 13206 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 33:

CGCTTTATCG TGGACGTGGT CAAGCCGAGA ATTTCATCAA GGAGATGAAG GAGGGATT	60
TTGGCGATAA AACGGATAGT TCAACCTTAA TCAAAAACGA AGTCGTATG ATGATGAGCT	120
GTATCGCCTA CAATCTCTAT CTTTTCTCA AACATCTAGC TGGAGGTGAC TTCCAAACTT	180
TAACAATCAA ACGCTTCCGC CATCTTTTC TTCACGTGGT GGGAAAATGT GTTCGAACAG	240
GACGCAAGCA GCTCCTCAA TTGTCTAGTC TCTATGCCCTA TTCCGAATTG TTTTCAGCAC	300
TTTATTCTAG GATTAGAAAA GTCAACCTGA ATCTTCCTGT TCCTTATGAA CCACCTAGAA	360
GAAAAGCGTC GTTAATGATG CATTAAGAA CAGTCGAGAT GAAAAAAATCG TGTGACGCAC	420
CAAGGGAGGA GTCTGCCCTT TTGAGGAAAT CTAGCGAGGA AAAACGATAC TGGAACAGCA	480
GAAAGTAAAA CTGACCTCAT GAGGAGGAAG AAAGTGGCTC ATGAGGTCA GGGTTTTGTA	540
AGTTACATCT AGTTGAGAGA GGTATGAATG ATTTGGGATT AATCATTTCT TGTTTTAAAT	600
CAGGAGAATA GTAACGATTT TTTCCTTTTG TGACGAACCTC TATTCCGTAA CGATCAATCA	660
ATTTAATCAT GTACCTAATA TTAGAATTGT TTATCCAAA TTATTTGAA AGCTTCTCTA	720
AGCTATATCC TTGTCTCTA AGTCATAGA TCTGAACCTT ATCATCATAA GTTAGTTCA	780
TAATAAAAAC ACCCCAAAG TTAGATTTT TCTGTCTAAC TTTTGGGGGG CAGTTCATTC	840
AACACCTGAT ACTATGCGTT TTTCTTATTT GAAATACTTT TTACTCAACC TCTTTATACT	900
CAATGAAAAT CAAAGTGCAA ACTAGAAAGC TAGCCTCAGG CTGCTAAAA CAGTGTGTTG	960

333

AGGTTGCAGA	TGGAAGCTGA	CGTGGTTGA	AGAGATTTTC	GAAGAGTATT	ACTTAATCTT	1020
CTTGATACTT	TGACTAAGAA	AAATCCTAC	AATCATCCCT	ACCATATTTT	GCATAAAATT	1080
CGGTAGAATT	TCTGGGAGGG	CTGCTGCCA	GCCATTCA	AAAGCAGAAC	CCAAGCGTA	1140
GCCTCCTACC	ATGCCAATAG	TTGCTAAAAT	AAGGCCTAAC	CACTGACTTT	TTCCCTTAAA	1200
TCCTGCGAAA	AAATCCTGCA	AGCCATGGTT	GACCAAGCTA	AAAGAACATCC	ACTGAGGGTA	1260
GCCTGATAAG	AGGTCAATCA	AGAAAATGTC	TAGTCCTCCG	ACTACCGCTC	CTTCACGACT	1320
ACCAAAGTAA	AAGCCGCAA	AGAAGACACC	AGCATCTAAA	AGAGTTAGAA	TTCCCTGTAGG	1380
TGTTGGGATT	TTTAAAGAAA	AAACCTAGAAC	CACAGAAAGG	GCGGTTAATA	GGGATACAAG	1440
GGCGATTTTA	GTTGTTTTG	TTTGCTTCAT	ATTGCTTAC	TCCATACTGA	TCTGCTTGTG	1500
CAATAGCAG	ATAAACGAAA	GCCTTAGAGC	TTTCTACTGC	TGGAAAAGT	TTATCACCTT	1560
TAACCAGGTG	ACTGGCAATG	CTAGAGsCAA	AGGTACAACs	TGCACCAAGCA	TTTTGGCCTT	1620
GGATAACTGG	ATTTCTAGG	ATAGTAAAGG	TCTGTCCATC	ATAAAAGACA	TCCACAGCCT	1680
TGTCCTGACT	AAGACGATTG	CCTCCCTTGA	TAATGACTG	GGCGCTCCTA	AATCATGCAA	1740
TTTCTGCGCT	GCAGTTTCA	TGTCTTCAA	GGTTTTAATT	TCCTGACCGG	ATAATAATTC	1800
TGCTTCTGGG	AGATTAGGCG	TAATCACACT	GACATAAGGG	AAAAAGCGAA	TCAACTCTTG	1860
GCAGAGCTCA	CTGACAGCTA	CATCATGCGT	TTCCCTGCAG	ACCAAGACAG	GATCCAACAC	1920
CACAGGTACT	CCTGGCGTT	GTTCATAAA	GTCCAAGGCC	TTCTCAGCCA	CGCTGACAGT	1980
AGGGAGAAGA	CCAATCTAA	TTCCCCAAA	TTCCACATCA	CGCAAGCTAT	CTAATTCTATG	2040
TTGAAAAATG	GTATCATCAG	TTGGAAAGAC	TTCAAATCCT	TTTCTGTCA	AGGCTGTCAA	2100
ACAAGTCACT	GCTACAAACC	CATGCAAGCC	GTTCAGGTA	TAGGTAGCCA	AATCAGCTGA	2160
CAGTCCACCA	CCACTAAAAAA	TATCATTTCC	AGAAAGTGCT	AAAATACGAT	TATTCTTCAT	2220
AACGAATCTC	CTTTAAATAC	AAACCATTG	GTGCTGCAGT	GGGACCTGCA	AGTTGCCTGT	2280
CCTTCCTCTC	CAAGATGAGA	TCAATCTGCT	CTACTGGCAT	GCGGTTGTTA	CCGATTTGA	2340
GAAGAGTCCC	CACCATATTG	CGAATCTGTT	TATACAAGAA	ACCATTCC	GAAAAGGTAA	2400
AGGTCAAAAA	TTGTCCTGTC	TCATCGACTA	TTAAACTAGC	TTCTGTGATG	GTGCGAACCT	2460
TATCCTCTAC	ACTAGTCCC	GAGGCTGTAA	AACCGGTAAA	ATCATGGTT	CCCTCTAGCT	2520
TTTGATTGCA	AATCTGCATT	CGTCCACAT	CGAGTGGGTA	GGGAAAGTGG	GTGGCATAGT	2580
GACGGCGCAT	CGGATTTTG	GGACGTCC	TATCCACAGT	AAACTCATAG	GTCTTGCTAT	2640
GCTTGGCATA	ACGGCAATGA	AAATCATCTG	CCACAAGCTC	AATCGAAATC	ACATCAATAT	2700

334	
CTTCAGGAGA CTGGGTATCC AAGGCAAAAC GGAGTTTCTC CTCATCCATC TGATAAGGCA	2760
GGTCAAAATG AATCACCTGT CCCAGGGCAT GAACCCCCT ATCTGTCCTA CCAGCACCCT	2820
GAACAGTAAT GGCTTGCCTT TTATTTAATC TGGTCAAGGT TTTTTCAATT TCTTCCTGAA	2880
CGCTACGCGC ATGAGGCTGG CGCTGAAAGC CAGCAAAGGC ATAACCATCA TAGGAAATAG	2940
TTGCTTTATA TCTCGTCATA GCCTCTATTT TATCAAGAAA TTAGTCTGTA AACAAAGGACC	3000
TAAACACAAAT ATTGTATGGG TATAAAAATC TCATACTCTT CGAAAATCTC TTCAAACACAC	3060
GTCAGTTCC ATCTGCAACC TCAACACACT ATTTTGAGCA ACCTGCGGCT AGCTTTCTAT	3120
AGTAGATTGA AATAAGATAT GAACAACCTCT ATTAGGAAAG TCACAAATTAT TTCTAGAAAT	3180
ATTTTAGCAG CTACAGCGTA CTATTCCAAA CTCACATCAAC TATAGTTGC TCTTTGATTT	3240
TCATTGAGTA TCAAAAGAAA AACTTAGGAA TCAATCCTAA GCTCTCTTCT GAAGTAGGTA	3300
CATGACAAAG ATAGAGATTA CAATCAACCA ACCTCCTAAG ATACTAAAGA CCAACATCCC	3360
ATTGTGAGTT AGTAAGCCAA TTGCACCTAG AACGAATGGG GTCGTAAAGG CTCCGAAACT	3420
ACAGCCTAAT ACAGCAAATG AAGTTGCTTG ATTGAGGAGT TTAGCTGGAA TTCGTTCAGA	3480
GACAAGTTGA AAGACCGTCG TCAAGACTAC ACTATAGGCA AATCCAGGCC GAACACTTCC	3540
TGCTACTACC ACCCACAAGG ATGAAGACAA GGCAATCACG ATTTGCCCA AGCCAAAGGT	3600
AATACCAGAC CAGAGGAGCA GTTTCTCTTT AAAGATAGAA ATCAAGAAAG AAAAACTCAC	3660
CCCAGCCACA ATCCCGATCA ACTGCATGAT ACTAAGAACAA AAACTAGATA ACTGGGCATC	3720
CCCCAATCCT CTTTCCACCA TCAAACATTGG AATACGGATG GTAATAGCTG TATTGGTACA	3780
AACTACAACG GCCGCTTCGA TAGCTAAGGT AAAATCAAG CCTTTCATTT CTCGAGTTAA	3840
ACGACTTGCT TCCTTCGCTC TTTTCTTGAC TTCTTTCTTT GATTTCCAT AAGGGACAAA	3900
GAGCAGATAA AGGGCAGCA CCAAAATCC AGCACTATAG GCTAGAAAGA TAGCTGTCCA	3960
ACCAAAGGCC ACAACTGAC CGACGGCCAA GGTAATGAGA GAAGCTCCAA CGACCTCTGC	4020
AGAACGCGGT AGCCCTAACCA TCTGAATTG CTTTTTCCT TGGTAGCGTT CACTGATAAT	4080
AGAAATGGCC TTGGCATTGA TCATCCCAAG ACCCAAACCA AAGAGAAGCC GTGTTCCAAA	4140
GACAAAGGGA TAGGCTTGGT ACCAGAAGGG AGCTGTACCG CTCACATGATA AAATCAGCAA	4200
GCCCAAACCA ATCTGTAAGC GCTCAGGAAA TATTTTTCTT AAGAAACCAT TTAGCAGTAA	4260
CATCATCATG ATTCCAAAGG AAGGCAAGCT CACCAAGAGC TCAATTGTT CCTTGAATA	4320
ACCCGTATAA TAGTCAAACA TGGCTGGTAG GGCACTCGAA ATGGAAAAGG AGGTAATCAA	4380
AACGAGGGAG AGAGCCAAA TGCTGGCCCG TTCTAAAAT TGTTTCATGA AATCTCTTTC	4440
TATATTTCTC TTAATCTTCT ACTTTTTGAA TAGTTATCAA ATAAGCAAGA AAAGAAGAAG	4500

CCTCATTGGT TTGTAGACTC CTTCTTAAAT TCGAAAATGA ATCCCTGTA TCTTATACTC	4560
AATGAAAATC AAAGAGCAAA CTAGGAAGCT AGCCGCAGGT TGTCAAAAC AGTGTGTTGA	4620
GGTTGCAGAT GGAAACTGAC GTGGTTGAA GAGATTTCG AAGAGTATTA GGATGACTTT	4680
CTCTTGATTT GCTTGATAAA GTAGAAAATA AATCCTGCTA CCATATAGGC AACAAAGATA	4740
ATCAGACACC ACTTAAACAC AACATTCCAA CCCTTGTTC AATTCAAAA GAAGTAAGGG	4800
AAAGGATTAT CCTTGGCATT TGGAAATATG AGTTTAGAA CCAAGCCATT AAAAGAGCA	4860
AACATCATAT ACAGAAAGGG TAAAATGGTC CACACTGCTG GATCCCAAAT CTTGTATTGA	4920
CCCTGTTTGT CAAAAAAGAG GGTATCCGCT AAAAACAGA TGGGAACGAT ATAGTGGCAA	4980
AGGAAATTCTT CTAGGGTATA GAAATTAGTC GCAATGGCG CCAAGAGGAA ATGGTAAATC	5040
ACACAGGTAA TCATGATACT CATGGTGACC CCACCTTTA AGCGCAAGAG ACTTGGCCTT	5100
TGCCAATTCTT CACCTACACG GCTCATACC TTTAGAAGAT AAAGGGTAAAT AATAGTTACC	5160
AAGAGGTTGG ACAGAACCGT GTAATAGAGA AGCATCCAA AACACCATG CTTAGTAATT	5220
TCAAGATAAA CTCCCGTAAA AGCCGCTAGA AACAGAAAGA TACGGCTATA AAATACAAGT	5280
TTATAGTGT TTAGACATGCT TAAATCTTCC TCACAAACTC TGATTTAAGT TTCATGGCAC	5340
CAAAACCATC AATCTTACAG TCGATATTGT GGTCGCCTTC TACGATGCGG ATATTTTCA	5400
CGCGCGTCCC TTGTTTCAAAT TCTTTGGCG CACCTTTTAC TTTCAGTCC TTGATGAGAG	5460
TTACTGTATC ACCATCAGCC AATTATTTTCC CGTTGGCATC GATAGCGACA AGACCTTCTT	5520
CTACTTCTGC AACTTCAGCA GGATTCCACT CATGAGCACA CTCTGGCAA ACCAGTAGGG	5580
CACCGTCTTC GTAGACATAC TCTGAGTTAC ATTTGGACA ATTTGGTAA TTGTTCATGG	5640
TTTCTCCTTA TCATCATTCA CTATTCTTG AAAATCAAAA TTTCTCGAAC AGCAACTATT	5700
ATACCCCTAAATC ATCAGCATTG TGACAAATTT AGAAAAAAAC CGATATCAAT CTATCGGCTT	5760
TTCTACATTT ACATTCTTTT TTCACTTCTG GCTTTGATTT TTTCAACTAC TTCTGAAATG	5820
TTCAAAACAG TTGTATCAAG GTAGACAGCA TCCCTCTGCTT GTTGGAGAG AGAAGTCTCA	5880
CGATGACTAT CCTTGTACTC ACCCGCAGCA ATTTCCCTTT TTAGGGTTTC AAGGTCTGTT	5940
TCAATTCCCT TGGCAATATT TTCTTGATAA CGACGCTCTG CTCTCTCATC AACAGAAGCT	6000
ACTAGGAAAA TTTCAATTC TGCTTGTGGC AATACAACAG TTCCAATATC GCGACCATCC	6060
ATGACAATCC CGCCTTGCTG GGCAATTCT TGTTGGAGAG AAACCAGTTT CTCACGCACT	6120
TGAGGAATTG CTGCAATAGC AGAAACATGA TTGGTCACTT CATTTCAGC GATAGGATGG	6180
GTAATATCCA CATCTCCTAC AAAAACAAAGC TGGTCTCCAG TTTCTGAACG TCCAAAGCTG	6240

336

ATTGGATGCT	GGTCCAACAA	GGCTAGAAGG	GCTTCGACTT	CTTCAACTCC	TAATTGGTTC	6300
TTAAGAGCCA	TATAGGTCGC	TGCACGATAC	ATAGCTCCTG	TATCAAGGTA	GGTGAATCCA	6360
AAATCCTTAG	CAATAATCTT	TGCGACCGTA	CTCTTACCGC	TGGAAGCAGG	ACCATCAATA	6420
GCAATTGAA	TTGTTTCAT	ATCGGCTCCT	ATTTTATTTT	TATAACATCA	CCTGGATTAG	6480
CAAACCAAGA	TCCTGTAGCC	ATGTGCCAG	GATTCAAGGC	CTCTAACTGA	GCAATGGAGA	6540
TTCCTGCACG	AGCGGCAATA	GCTGTTCCC	CTTCTCCTGC	GAGAACTTTA	ATCGTTCTT	6600
CAGGATTAGC	AGCTCTTCT	GAACTACTAG	AAGTAGATTC	TGGCTCTGAA	CTCTGCTCAG	6660
GCTGAGAACT	ACTTGAAGAT	GAGATTTGTA	CTACACTGGC	ATCAGAAATCA	TGAAAGCCTT	6720
TTAAGGCTGC	TGTGCGATTA	CTCCCCCCC	ATGATAGATA	GATGAGAAC	ATGACCATCA	6780
CCACCACAAAT	TACAAAGAAA	ATACTAGCTA	GGATCGTCAA	AATACGATTA	GCCATCCTAT	6840
CAGCCCCCTCC	GTGGTTTCGA	TGCCGACGCT	CTGCTCTTGA	TTCTTCTTGA	TCATAGATAT	6900
CTTCTTGCCA	CGGTTCTTT	GCCATACCTT	ACTCCTTGTT	TTTTTTTACT	TTTCTTATTA	6960
CAATATAAAT	ATGAACATGA	AAATCACACT	TATACCTGAA	CGATGTATCG	CCTGTGGGCT	7020
TTGCCAAACT	TATTCGTATT	TATTTGATTA	CCACGATAAT	GGAATCGTGC	TTTTTTACGA	7080
TGACCCCTGAC	CAACTGGAAA	AAGAAATTTC	TCCTAGTCAG	GATATCTTAG	AGGCTGTTAA	7140
AAATTGCCCA	ACTCGCGCCC	TGATTGGAAA	CCAGGAAGCC	AAATCAATG	GCGATAATCC	7200
ACTCCCTCTA	GTTTAGCACA	TTTCCATGTA	AAATTATAGT	CTTTTCACTT	TATTTTTTTC	7260
TGTAAAATCA	GGAAAGGTCA	TTTTTCTTT	GATAAGATAA	AGTGGTCTTT	TTTTAGTCTC	7320
TAAATAAATC	TTACTGATAT	ACTTGGCGAG	AATCCAATG	GTCAAGAGTT	GAATGCCCTCC	7380
AAGAAAGAGA	ATAACAGCCA	TCAGAGAGGT	CCAACCAGAT	GTCGGATTGC	CCAAATGAG	7440
GGTCCGAACC	ACAACAAAAA	AGGTCATCG	CAGAGAAAGA	AAACAAGATA	GGAGACCAGC	7500
TACAAAGGCT	ATAATCAAGG	GAAAATCTGA	AAAATTAATA	ATCCCTCAA	TGGAGTAGAA	7560
AAAGAGTTGC	CTAAAACCTCC	AACTTGTCTT	GCCAGCCTGC	CTTTCGACAT	TTGGATAGTC	7620
CAAATAGTAG	TTTTGAAAC	CCACCCAGGC	GAAGAGCCCC	TTTGAAAAAC	GATTGGACTC	7680
GGTCAAGCTT	AAAATGGCAT	CGACTACAGA	CCTTCTCATC	ATACGAAAT	CACGGACACC	7740
CGACGGCAGA	GCTACTGGGG	TGATTTTTTG	CATGAGGGCGA	TAAAAGAGAA	CAGCACAGAA	7800
ACTGCGAAAG	AAGGGTTCTC	CCTCCCGACT	AGTTCTCCGT	GTCCCAACCG	AGTCCAAGTC	7860
TACATTTTG	TCTAATACAT	TTTCATCTC	AAACAACATA	CTAGGAGGAT	CTTGGAGGTC	7920
TGCATCCATC	ACCACCCACCA	AATCTCCTGT	CGCATATTGC	AAGCCTGCAT	AAAGGGCTGC	7980
TTCTTGCCA	AAATTCGAG	AGAAAGAAAT	ATAATGGACT	GCCGGATTTT	GCTCCCGATA	8040

GGCCTTAAG AGTCCAAGG TCCCACACT TGATCCATCA TCGACAAAGA CATACTCGAT	8100
TTCTGTTCC AAATCTGGAA GTAAAGCTTC CAGAGCCTGA TAAAAAAGAG GAAGTACTTC	8160
CTCTCGTT AAACAAGGGA CGATGATTGA AATCATCATC TTAGTCTTC AATCCATTG	8220
GATGCTTGCT TTGCCAACGC CATGCGTCTT CACACATTG GGTGATGTGC AGTTCTGCTT	8280
CCCAACCGAG TTCTGCTTA GCTTTGCCG GGTCTGAGTA GCAGGCAGCG ATATCACCTG	8340
GGCGACGTTT TACGATGCGG TAAGGAATAG GACGGCCAC CGCTTTTCC ATGTTTGGA	8400
TAATTCAG AACTGAGTAA CCTTTACAG TTCCAAGGTT ATAAACGTT AGTCCTGAAC	8460
CTTTTGGAT TTTTTCAAA GCTGCAACGT GACCTTACG CAAATCGACA ACCTGGATAT	8520
AGTCACGAAC ACCTGTTCCA TCTCCCGTAT CGTAATCGTC TCCAAACACT TGCACTTGCT	8580
CTAATTTCC AACGGCTACT TGAGTCACAT ATGGCAAGAG ATTGTTGGA ATACCGTTG	8640
GATTTCTCC CAAATCACCA CTCTCATGGG CTCCGATTGG GTTAAAGTAA CGAAGCAAGA	8700
CAACATTCGA TTCTGAGTCT GCTTTGTAAA TATCAGTCAA AATTTCTCT AGCATGAGCT	8760
TAGTACGACC GTATGGTTG GTCACTGAAA GTGGGAAATC TTCCAAGATG GGCACGTGTT	8820
GCGGATCCCC GTAAACTGTC GCAGAAGAAC TGAAGATGAT GTTTTACAG TTGTTTCTT	8880
CCATGGCTTT CAAAAGGCTG ACAGTTCCAG CGATATTGTT GTCATAGTAG GCAAGAGGGA	8940
TACGTGTTGA TTCCCAACAA GCCTCAACAC CAGCAAAGTG AATGACACCA GTGGTTCTT	9000
CCTGCTTGAA AATATCTCTG AGGGTATCTG TGTCAACGAA ATCTGCCTCA TAGAAAGGAA	9060
TCTCAACTCC TGTGATTCTC TCAACAACTT CTAACACTT ACGATTGCTA TTGACAAGAT	9120
TATCCACAC AACAACTTGA TGACCTGCTT GGATCAATTC AATAACAGTG TGGTTCCAA	9180
TAAAACCGGC ACCACCACTT ACCAAAATCT TTTCTTGAT CTTTTTCTT CGATTCTCAG	9240
ATTATTTTTT CTTATTTTAC CATTGGAC AGGGATGTC ATTGCCATC CTAACACTACC	9300
TGATAAAATT TCAGTAAAT GCTTATACTC TTCGAAAATC CAATTCAAC TACGTCAACG	9360
TCGCCTTGCC ATGGGTATGG TTACTGACTT CGTCAGTTCT ATCCACAACC TCAAAACAGT	9420
GTTTGAGCT GACTTCGTCA GTTCTATCCA CAACCTCAAA GCAGTGCTT GAGTAACCCG	9480
CGGCTAGTT CCTAGTTGT TCTTGATTT TTATTGAGTA TTATTGCTT TTTACTCGTT	9540
TGACATAGTT TTCAATTGGG TAATTTAGAG GGTCCAAGGT CAACTCCTTG TCTTGGATCA	9600
GTTGGGCTAG ATGGTAACCA ATGATAGGAC CAGTTGTGAG GCCTGATGAA CCTAGTCCAC	9660
TGGCTGCATA GACACCAAGTT AAGTCAGGCA CCTGCCAAA GAAAGGAGAG AAATCACTGG	9720
TGTAGGCACG GATTCCAACAA CGCTCAGATT TTGAAGTAGC TTCAGCCAAA ATCAGATAGT	9780

338	
GAGTCAAGGT	9840
GGCCTCCTCC	
ATTTGTTGGA	
GCAAGGTTTC	
ATCTACCGTC	
AAATCAAATC	
CCATGTCATT	9900
TCCTGGGTA	
GGCCCTAAGG	
ATAATTCccc	
ACCTGCAAAG	
GGAATCAAAT	
CCCACTCCCC	9960
TTCTGGCATG	
ACAAACAGGGT	
AATCTTCCAT	
GTCTTGGCA	
AGCTGATAAT	
CTCGTAGTTG	10020
TCCTTTTGA	
GGACGGACAT	
CCACTTCATA	
ACCTAAAGGC	
TCTAACATGT	
CCCCCAACCA	10080
AGCTCCCGTC	
GCCAAAATAA	
CCTGCTCAA	
CTCCTCTTC	
CCAATCTGGT	
AGCCTGATGC	10140
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AGAGTCACTT	
TTTCCTTGAC	
CAGCTTGACA	
TGACTGACTT	
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TCCACCAGAA	
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ATTCAATTAGC	
TGAGGCTTGG	
TTCAGAATGG	
CTAATTGCC	10320
TATCAAGGGA	
GATTCTTCTC	
TGCGCTGGAG	
GGCCAGTTGA	
TAAGATTCTT	
CCAAATTGGA	10380
TTCATCCTTT	
TTCAAGAGAA	
AGACTCCCGA	
ACGCTGGTAA	
AAAGTCGATTT	
CTTGTCTGA	10440
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AAAATCAGCC	
CCCAAGCGCG	
CCATCTTGTA	10500
CCAGGCTTTA	
TTACGGCGTT	
TGGAAAACCA	
AGGACTGATA	
ATTCCCTGCTG	
CGGCCTTGGT	10560
GGCTTGACCT	
TGCTCATGGT	
CAAAAACGGT	
CACCTCTAGG	
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TCGAGAGGTA	10620
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ATAGTGACCT	
GGGTCCAATG	
CAAGCAACCC	
ATCAGATAGC	
ATATCCTGAG	10860
CAGTATGGTA	
GTAGATATCA	
CCAGTGATAT	
AGACATCTGC	
CCCCTTTGCC	
AAAGCATCCT	10920
TATAGAAAGA	
CTGCCCGCTT	
CCACCACAAA	
TTGCTACTCT	
TGAAATAGGC	
TTCTGCAAAT	10980
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ATAATGCACC	
ATTGGAAGGC	
TATCTAGGTC	
AAAGACTTGC	
TTGACCTGTT	11040
GGGCCAATTC	
CCAAAATGTC	
TGAGGCTGAA	
TATTCCCAAT	
ACGTCCAATT	
CCACGTTCTG	11100
GACCTGTTTC	
CTGCGAGATA	
GTCGCTCCT	
CGATTCCTAG	
CATCTGACAA	
AACCAGTCAT	11160
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GATGAAATTC	
TGATTTGGCG	
GACGGCTGGC	
AAGCAAGTCC	
TTGATAGGAC	11280
GAAAGATAGG	
CGCGTGTGTTG	
ACGATAATCA	
AGTCCACACC	
CTTTTCAATG	
GCCTCTGCCA	11340
CTGCTCTTC	
ACGAATATCG	
AGGGCAACCA	
TGACCCCTTG	
GATACCCCTG	
TCTAAAGTGC	11400
CAATTGCGAG	
ACCACGGCTG	
TCTCCCTCCA	
TAGAAAATTC	
CTGAGGGCAA	
AAGGCTTCAT	11460
AAGCTTGGAT	
CACTTCACCT	
GCTAACATGG	
AGCACCTCCT	
TGATAGCTTG	
AATCTTATCT	11520
ACTAGAACCT	
GACGTTCTTC	
CAGATTTTTT	
TCTGGGATTT	
GTCCGAGGGC	
GAACCTCTAGC	11580
TTCTCAGCTT	
CTTTTGCCCA	
TTTTTGGACA	
AATACTGGAC	
TGACTTCTTT	

GGACAAAGAAG GGACCAAAGC GAACATCACT GGCTGATAGC TTCAATTGTC CTGCTTCCAC	11640
CACCAAAATC TCATAAAACT TTCCAGCTTC TTCTAAGATG CTTTCTGCTA CAATCTGGAA	11700
TCCATGATCC TGTAGCCAGA TACGCAAGTC GTCTTCACGA TTATTGGGCT GGAGGATCAA	11760
ACGCTCTACA TTAGCTAACT TCCCCAAACC TTCTTCTAAA ATCCTAGCAA TCAAACGACC	11820
ACCCATGCCA GCAATGGTAA TGACAGACAC TTGGTCAGTC TCTTCAAAG CTGCCAAGCC	11880
ATTGGCTAAA CGGACTTGGA TTTTCTCCTT TAGGCCGTGA GCCTCAACAT TTTTAACCGC	11940
AGACTGATAG GGACCTTCCA CCACCTCACC TGCAATAGCG CTTTGATTT GGCTCTCTC	12000
AACCAACTCG ATAGGCAGAT AAGCATGGTC ACTTCCCACA TCTAGTAAA TAGCCCCCTG	12060
TGACACAAAG GAAGCTACCA ATTCTAATCT CTTTGAAATC ATCTTCTCTC ACTTTCCAAA	12120
ACTCTATTAC CTCTTATTAT ACCACATTTC AATCTTCAAC TTCCCAGTAA TATAAGCACC	12180
TCTGGCGAAA GAAGTTCAA TGTCTAAAG TAATAAGTGA ATCCAATTGA AAGATTTAA	12240
ACAATTGCA AAAATGTCAA AAAATAAAA ATAAACAGTT TATTCAGAAA ATTCTTGACA	12300
TATAAAAACA CATGGTAGAA TATAATTAGA AAGTTAGAAA AAATAAAAGT TTGACTAAAA	12360
TTTGTATTTG AAGGTGGTGT TCAGATAAGA AATTTAGTCA GACGAACCAC GAATTTGCTC	12420
TATGCTTTCT GGAATTATTC ATAACAGGAG GATACAGTCA TGGAACAAAC ATTGTTGAA	12480
TTAGAACTAC TTCCAGAGGA AGATATCATT GTCACAGGTC TCCCTAAGTA TTGTTCTTT	12540
ACTTGTAAATTTA TTACAGGTCT CTAGTTATAT TTTATATAAA ATAAGTAGCT TTACTTACGG	12600
AATAGGCTAG TGCTGTGTCT CTAGCCTATT TTAATAATTA GGAGTTGTT ATGGATTAT	12660
TAGAGAAAGA ATGTTAAAAA TGTGATAAAA ATTTCCAACA GGGTGATATT TGGAATTACT	12720
ATTATTTATC AGATAAGATG CCTGCACAAG GGTGGAAAAT ACACATAAGC TCCCAATAA	12780
AAGACGCTGT AAATATTTT AAGATTGTGT ATAAACTATC CCAACTAAAT AATTGTAGCT	12840
TTAAAGTTGT TAAAAATTAA GAGGAATTAA AAAAATTAA TTCCCCTAGG GAAATGAGCC	12900
CTACTGCTAA CAAATTATA ACTCTATATC CTAAGTCAGA ATCTGAAGCT AAGAGTATGA	12960
TTTGTAAATCT TACGAATAGA CTGTCAGAAT TTAAGGCTCC AAAAATACTA TCTGACTATC	13020
AATGTGGAAT GCATTCTCCA GTTCATTATA GATATGGGGC TTTTTAAAAA AAACAAGCTT	13080
ATGATGAAAA AAATAAAAAT GTCACTATT TATTGCTAGA TGAAAAAGG AAGAACTATG	13140
TAGAAAGATAA GAGACAAAAT TTCCCTAGTC TTCCTAGCTG GAAAATGGAT TTATTTCAAG	13200
AAGAAG	13206

(2) INFORMATION FOR SEQ ID NO: 34:

340

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 13104 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 34:

CCGGATCCAG CGAAAAATAT GCTCTTGAT GCTGTAAGTG GTCAAAAAGA TGCTAAAACA	60
GCTGCTAACG ATGCTGTAAC ATTGATCAA GAAACAATCA AACAAAAATT TGGTGAATAA	120
AAAATTTGTT CAAGGGGGGT GGAAATCAA TCCCCCTTG AATTATCAA TAGAGACACA	180
AATAATTTAG CTTTCTTATA AAAAAGTAGT ATCCTATGAA AGGAGTTAAT ATGGAAAAGC	240
AACAAACCTAG TAAAGCAGCC CTGCTGTCTA TCATTCCTGG GTTAGGACAG ATTTACAATA	300
AACAAAAAGC CAAAGGTTTT ATCTTCCTTG GTGTAACCAT CGTATTTGTC CTTTACTTCC	360
TAGCACTTGC AACCCCTGAA TTGAGCAACC TCATCACTCT TGGTACAAA CCAGGTCGTG	420
ATAATTCCCT CTTTATGCTG ATTCGTGGTG CCTTCCATCT AATCTTTGTA ATCGTTTATG	480
TACTCTTTA TTTCTCAAAT ATCAAAGATG CACATACGAT TGCAAAACGC ATTAACAATG	540
GAATTCCAGT TCCACGCACA CTCAAAGACA TGATCAAAGG GATTTATGAA AATGGCTTCC	600
CTTACCTCTT GATCATTCCA TCTTATGTTG CCATGACCTT CGCGATTATC TTCCCAGTTA	660
TCGTAACCTT GATGATCGCC TTTACCAACT ACGACTTCCA ACACCTGCCA CCAAACAAGT	720
TGTTGGACTG GGTTGGTTTG ACCAACTTTA CAAACATTG GAGCTTGAGT ACCTTCCGTT	780
CTGCCCTTGG TTCTGTTCTT TCTTGGACTA TCATTTGGC TTTGGCAGCT TCTACTTTAC	840
AAATCGTAAT TGGTATCTTC ACAGCTATCA TTGCCAACCA ACCATTATC AAAGGAAAAC	900
GTATCTTGG TGTTATTTTC CTTCTTCCTT GGGCTGTCCC AGCCTTCATC ACTATCTGA	960
CATTCTCAA CATGTTAAC GATAGTGTGCG GTGCTATCAA CACTCAAGTA TTGCCAATCT	1020
TGGCTAAATT CCTTCTTTTC CTTGATGGAG CTCTTATTCC TTGGAAAACA GACCAACIT	1080
GGACTAAGAT TGCCCTGATT ATGATGCAAG GTGGCTCGG ATTCCCATAC ATCTACGTT	1140
TGACCTTGGG TATCTTGCAA TCTATTCTA ACGACCTTTA CGAACGAGCT TATATTGACG	1200
GTGCCAACGC TTGGCAAAA TTCCGCAACA TCACCTTCCC AATGATTTTG GCTGTTGGG	1260
CACCTACTTT GATTAGCCAA TACACCTTCA ACTTTAACAA CTTCTCTATC ATGTACCTCT	1320
TCAATGGTGG AGGACCTGGT AGTGTGGAG GTGGAGCTGG TTCAACCGAT ATCTTGATCT	1380
CATGGATCTA CCGTTGACA ACAGGTACAT CTCCTCAATA CTCATGGCG GCAGCTGTTA	1440
CCTTGATTAT CTCTATCATT GTCATCTCAA TCTCTATGAT CGCATTCAAG AAACTACACG	1500

CATTTGATAT GGAGGACGTC	TAAGATGAAT AACTCAATTAA	AACTCAAACG	TAGACTGACT	1560	
CAAAGCCTTA CTTACCTTTA	CCTGATTGGT	CTATCAATTG	TAATTATCTA	TCCACTGTTG	1620
ATTACCATTA TGTCAGCCTT	AAAGCAGGT	AAACGCTCAG	CCTTTAAACT	AGATACTAAT	1680
ATCGACCTCA ATTTTGATAA	CTTTAAAGGC	CTCTTCACTG	AAACCTTGTA	CGGTACTTGG	1740
TACCTCAACA CTTTGATTAT	CGCCTTAATT	ACCATGGCTG	TTCAAACAAAG	TATCATCGTA	1800
CTTGCTGGTT ATGCTTACAG	CCGTTACAAC	TTCTTGGCTC	GTAAACAAAG	TTTGGTCTTC	1860
TTCTTGATCA TCCAAATGGT	GCCAACATAG	GCCGCTTGA	CAGCCTTCTT	CGTTATGGCG	1920
CTTATGTTGA ACGCCCTAA	CCACAACTGG	TCCTCATCT	TCCTCTACGT	TGGTGGTGGT	1980
ATCCCAGATGA ATGCTTGGCT	CATGAAAGGC	TACTTCGATA	CAGTGCCAAT	GTCTTTAGAC	2040
GAATCTGCAA AACTAGACGG	TGCAGGACAC	TTCCGCCGCT	TCTGGCAAAT	TGTTCTACCA	2100
CTTGTGCGC CAATGGTGC	CGTACAAGCT	CTCTGGCCT	TCATGGGACC	TTTCGGGGAC	2160
TACATCCTCT CTAGTTTCTT	GCTTCGTGAG	AAAGAACTACT	TTACTGTTGC	CGTAGGTCCTC	2220
CAAACCTTCG TTAAACAATGC	GAAGAACTTG	AGAGATTGCC	ACTTCTCAGC	AGGTGCTATC	2280
CTCATCGCCC TTCCAATCTG	TATTCTCTTC	TCCTCCCTAC	AAAAGAACTT	TGTTTCAGGA	2340
CTTACAAGTG GTGGCGACAA	GGGATAATT	ATCCCCGCCA	CCCTTTTCA	TTTTATACTC	2400
TTCGAAAATC TCTTCAAACC	ACGTCAGCTT	TATCTCCAAC	CTCAAAGTTG	TGCTTTGAGC	2460
AACCTGTGGC TAGTTTGAC	TTTGATTTC	ATTGATTATT	AGCAATTGTC	ACTGTAAATA	2520
ATATCCTTGT AGCAAGCAAT	TTTTCTCCTA	GACTTGAAAT	AAAGCGCATT	TCTCTATATA	2580
ATAATACTCA TATAGAAAAC	ACCTTTAGA	AAGATACCTA	TGCTTCCATA	TCCATTTCC	2640
TATTTTTCAA GTATTTGGGG	GGTTCGTAAG	CCCCTGTCCA	AACGTTTCGA	GCTCAACTGG	2700
TTTCAACTTC TCTTACCAAG	TATCTCCCTT	ATCAGCTTGT	CTATGGTACC	CATTGCTATC	2760
CAAACAGCT CCCAGGAGAC	CTATCCGCTA	GAAGTTTTA	TCGATAATGT	CTATGAACCT	2820
CTGACAGATA AGGTTGTCCA	GGATCTCTCT	GAACATGCTA	CAATTGTCGA	TGGCACATTA	2880
ACTTTACTG GAACAGCTAG	TCAAGCCCT	TCTGTTGTGA	TTGGTCCAAG	TCAAATCAAG	2940
GAATTACCTA AGGACTTGCA	ACTGCATTTC	GATACAAATG	AGCTAGTCAT	CAGCAAGGAA	3000
AGCAAGGAAC TGACCCGCAT	CTCTTACCGA	GCCATTCCAGA	CTGAGAGTTT	CAAAAGCAA	3060
GACAGCTTGA CCCAAGCAAT	TTCTAAAGAC	TGGTACCAAC	AAAATCGTGT	CTATATCAGC	3120
CTCTTCCCTAG TTCTCGGTGC	GAGCTTCCCTC	TTTGGTTTGA	ATTTCCTTAT	CGTCTCTCTT	3180
GGAGCTAGCT TTCTCCTTTA	TATCACCAAA	AGATCACGCC	TCTTTTCATT	TAATACCTT	3240

342						
AAAGAGTGCT	ACCATTTAT	CTTGAACGT	TTAGGATTGC	CGACTCTGAT	TACACTTATT	3300
TTGGGATTAT	TTGGCCAAA	TATGACAACC	CTGATTACTG	TACAAAATAT	TCTTTTGTT	3360
CTGTATCTGG	TCACTATCTT	TTATAAAACA	CATTTCCGTG	ATCCAATTAA	CCATAAATAG	3420
GAGATTTTA	TGCCCCTTAC	GATTAAAGAC	GTGGCCAAGG	CTGCTGGTGT	TTCGCCCTTCA	3480
ACCGTAACCC	GTGTTATTCA	AAATAAATCA	ACCATTAGCG	ACGAAACAAA	AAAACGTGTT	3540
CGCAAAGCTA	TGAAGGAAC	CAACTACCAC	CCAAACCTCA	ACGCTCGTAG	CTTGGTAAGC	3600
AGCTATACTC	AGGTTATCGG	ATTAGTTCTT	CCTGATGACT	CAGACGCCTT	CTACCAGAAAT	3660
CCTTTCTTTC	CATCGGTTCT	ACGTGGCATC	TCTCAAGTCG	CATCTGAAAA	CCACTATGCC	3720
ATTCAGATAG	CAACAGGGAA	AGATGAGAAG	GAGCGTCTCA	ACGCTATTTC	ACAAATGGTC	3780
TACGGCAAGC	GTGTAGATGG	GCTAATTTTT	CTCTATGCC	AAGAAGAAGA	CCCTCTCGTA	3840
AAACTCGTCG	CAGAAGAAC	GTTCCTCTTC	CTTATCTTAG	GTAAATCTCT	ATCTCCTTTC	3900
ATCCCACCTG	TCGACAAACGA	CAATGTTCAA	GCTGGTTTTG	ATGCGACTGA	ATATTCATC	3960
AAAAAAGGCT	GCAAACGCAT	TGCCTTATC	GGAGGAAGTA	AAAAGCTCTT	CGTGACCAAA	4020
GACCGTTAA	CAGGCTATGA	ACAGGCGCTT	AAACATTACA	AACTTACAC	TGACAACAAAT	4080
CGCATCTACT	TTGCCGACGA	GTTCTGGAA	AAAAAGGGCT	ATAAATTTAG	CAAGCGATTA	4140
TTCAAGCACG	ATCCACAAAT	TGATGCTATC	ATCACAAACG	ATAGCCTCCT	AGCTGAAGGT	4200
GTGGTAACT	ATATTGCCAA	ACACCAAGCTG	GATGTCCTG	TTCTCAGCTT	TGACTCGGTT	4260
AAATCCAAGC	TCAACTTGGC	AGCCTATGTC	GATATCAATA	GTTTAGAGCT	TGGTCGTGTT	4320
TCCCTTGAAA	CTATTCTCCA	GATTATTAAT	GATAATAAAA	ACAATAAAC	AATTTGTTAC	4380
CGTCAATTGA	TCGCCAACAA	AATTATCGAA	AAATAAGAGA	CTGGGCAAAA	AGTCGTTAAA	4440
AGCAAAACG	CATACTATCA	GGTATTGAAA	AAACTTGATA	CTATGCGTTT	TATTGTGGGA	4500
AGATTTACTT	CCTTTCTAC	TGAAATTGAG	TCTTTCCCA	AGATCTTTT	ATACTCAATG	4560
AAAATCAAAG	TGCAAACTAG	GAAGCTAGCC	GCAGGTTGCT	CAAAACACTG	TTTGAGGTT	4620
GTAGATGAAA	CTGACGAAGT	CAGTAACCAT	ACCTACGGCA	AGGTGAAGCT	GACGTGGTT	4680
GAAGAGATTT	TCGAAGAGTA	TTAATCACTA	ATTATCTATC	TCAACAAATC	TTCTAGAAT	4740
ATGAACATTT	TCCGAGACAG	AGACAAAGGA	GCTTGGATCC	ACTTGTGTCA	TAATCTGTTT	4800
AAATTCAATTA	AACTCTGCAC	GTGTAATGAC	AGTGATTAAA	ACTGCCCTTC	TCTCGTGATT	4860
ATAGGTTCCCT	TCTGCATCGT	GGATCATGGT	TGCTCCGGCGG	TGCAATTTTT	TATGGATTTT	4920
TTCAATTACC	TTCTCTGGAT	GATTTGTCAC	AATCATGGCC	TGCATACGCT	TTTGCTTAGT	4980
AAAGACTGCG	TCTGTCACAC	GGCTAGAGAC	AAAGATGGTA	ATCATAGAAAT	AAAGAGCGTA	5040

TTTCCAACCA AAGGTCAAC	CTGCTATCG CATGATAGTT	CCATTTACCA AGAAAGAAAT	5100
ACTACCGACA TTCTTACCG	TTTCTTACG AATAGTCAGG	CTGACGATAT CCGTCCCACC	5160
ACTGGAGATA TTGTTTCGAA	GAGCAAAACC AATCCCCAA	CCCATAACAA CACCCCCAAA	5220
AAGGGAATTG ATAATGGGAT	CCTCTGTCAA GGTTGCCACA	GGGACAAACT GGATAAAGAA	5280
GGAACTCATA GATACCGTGA	TAAAGGTAAA GACGGTGAAC	TTATGGCCAA TCTGATACCA	5340
AGCTAAGACC ATCAAAGGA	AGTTAATGGC	CTAGAAGCTT AGCGAAATCG	5400
ACCAAACAG TGATTACTCA	AGGCAGAGAT	AATCTGTGCC AGACCTGTTG	5460
ATACACATGC CCTGGTTGGA	AAAAGAAATT	AACTGCTACT GCTGATAAAA	5520
CAGAGAGGCC GAAATCTCT	CATCATACTT	TTCTCGAGAG ATACTTGTAA	5580
AATTTTATC TGATAAGCAA	AGCGGCCAG	ATAATAGGCC CACCGCTTAA	5640
TTGTTTCATC TTCTCTACT	TGTAAGCTGA	GTCCTCTAG TTGTTTGAGA	5700
ATGGAGCTTG TGTCATTGGG	TCAGTTGCCT	TGTTGTTCTT AGGAAAGGCA	5760
GGATATTTTC TTCTCCAGCA	AGCAACATGA	CAAACGGTC AAGCCCGATA	5820
CGTGTGGTGG GAAACCATAG	TCCATGGCTT	CAAGAAGGA ACCAAACTGG	5880
CTTCAGTTGA GAAACCAAGA	GCCTTGAACA	TGCGTTCTTG AAGGTCTTT	5940
GAAGGCTACC ACCACCAAGC	TCATAACCCT	TCAAGACGAT ATCGTAAGCA	6000
CCTTAGCCAA ATCACCTCT	AATTCATGAG	CACTCTCTTC CTGTGGAAGT	6060
GGTGGCGCT CATGTAGCGG	CCTCTTCTT	CAGACCATTC AAACATCGGC	6120
CCCAAAGGAA GTTGAACCTA	TCATTATCAA	TCAAGCCAAG CTCTTAGCA	6180
GAAGGGCACC CAGTGTGCA	TTAGCCACTT	CAAGCGTATC CGCCACAAAG	6240
CCTTATCTTC AAGACAAGC	GCTGTTGTCA	ATTCTTCTTG GATACCGTC	6300
CAACTGGTCC GTTTAATTCT	CCATCAACCA	CCTTGACCCA AGCAAGACCT	6360
ACTGTTGGC TACTCCGTC	ATCTTGTGCA	TGTCTTTACG TGAATAGTTG	6420
CTGTGACCAAC AATCGTTTT	ACAGCAGGTG	CTTCTGAAAAA GACTTTAAAG	6480
GGACCACTTC TGTCAAGTCC	TGAAGCAACA	TGTCAAAACG AGTATCTGGC	6540
CGTAAAGAGC CATACCATCA	TCGTATTTCA	TACGAGGGAA TGGTAGCGTT	6600
CTTTGTTTC CTTCATCAGC	CGCGCGATCA	AGCTTTCTGT AATATCTGG	6660
CAGTAAGGAA GGACGTTTCC	AAGTCGACCT	GAGTAAATTC AGGCTGGCGG	6720
AGTCCTCGTC ACGGAAACAT	TTAACGATTT	GGTAGTAACG GTCAAAACCA	6780

344

AGAGCTGTTT CGTGATTTGAG GGACTTGAG GAAGAGCGTA AAAATGCCCT TTATTAACAC	6840
GAGACGGCAC TAAATAATCA CGCGCCCTT CAGGCGTTGA CTTAGAAAGG AATGGTGTCT	6900
CCACGTCGAT AAACCTCAAAC TCATCCAAGT ACTTGCGGAT AGAGTGGTC ACCTTGGCAC	6960
GAAGTTAACG ATTTCCAAC ATTTCTGGAC GACGAAGGTC AAGGTAACGG TAACGCAAAC	7020
GTGTATCGTC ATTTGCCTCA ATGCCATCCT TAATCTAAA TGGTGTGTC TTAGCTGTGT	7080
TAAGCACAAT AAGAGCTGTC ACGTTAACCT CAACCGCACC AGTTGGCAAC TTATCATTGG	7140
CTTGTACGC GCAGCGACCT GACCAGTCAC CTCATAACA AATTGCTAC GAAGGCTTTC	7200
AGCTGTTGCC ATAACCTCTG CAGATACTTT TTCAGGGTTG ATAACCAAAC GCATGATTCC	7260
TTCACGGTCA CGAAGATCGA TAAAGATCAA ACCACCAAGG TCACGACGAC GGCAACCCA	7320
TCCCTTCAAG GTTATTTCTT GTCCGATGTG TTCCCTACGA ACACGACCAG CATACTACT	7380
ACGTTCTATT ATTTCTCTCC TCTTTTATTC TGTTACTATT TTACCATAAA AGCCAGCTC	7440
TTCATGAAAA TCATCAGAAA AGTTGCCAG TCTTTAAAAG TCAGGTGAAA GCCCTAAAAA	7500
TTAGGCCTAA TACTCTTCGA AAATCTCTTC AAACACGTC AGCGTCGCCT TACCGTATGT	7560
ATGGTTACTG ACTTCGTCAG TTTCATCTAC AACCTCAAA CCATGTTTG AGCTGACTTC	7620
GTCAGTTCTA TCCACAAACCT CAAACAGTG TTTTGAGCAA CCTGCGGCTA GCTTCCTAGT	7680
TTGCTCTTG ATTTTCATTG AGTATAATAC AAAAATCCGA TGAACCTCAC CGGACTCTTT	7740
TATTTGAAT TTTTGCCCTGC TTTACGCTTT TCAGCGATT TCAGCTGCCCT TCGAGGCAAG	7800
ACAATTCCG TTATGTAAGC CGTCCCAAAA CGCAGTACAC CTGCAATAGG AGCAAAGACA	7860
ACTGCTAGAT AGTTATAGAA GAAATCGCCT TTGAAGGCAT AAGCTAGGCC TCCAATGATG	7920
AAAAATAGAA CGACTGCCTG AATCACTGCT AATAAAATTA CTCGTTTCAT GTGACCTCCT	7980
GACTCTATTA TAGCATGAGA ATCATCAAA AGCCGACTAA ATTATTCAA GCGTGAAGAG	8040
AAATACTGTA GACCAGACCT TTTCTGCTAA TGTAAGCCAA ACCCAAACCA AAACCAAGGC	8100
AAAAATAGAC AAAAATGTG TGACATCAC CTGGAAAATG AATCAAGGCA AATAGAAGAC	8160
TAGATACCAG AAGAAAATC AGGGTTCGTT TACTATTGTC CTGCTTAGGA AAGAGATAGC	8220
GTGCTAACAT CCCTCTAAA ACAATCTCTT CCGTCAAAGG AGCAAAATA ACCACAGCAA	8280
AGAATGAGAA AAGTGGTTGA GACAAGGTCA AGTCTGTCGC TATTTGCTGA TTTACTGAAG	8340
GATCATCTGG CAAGAAGAAT TGAACGACCA GAGATAAGAA CCAAACCAAG ACAGGAAGCC	8400
AAATAAAATCG ATTAAGCCG CTCTCTCAA TATGAACAGG AGCCTCTGA TACCATTTGT	8460
AAATGCCGTA CACATATACT CCAGCCAAGG CCACATAGAG TAGAGTAACA GCATAGGGTG	8520
AAGGCCTAA AGCAAGCGAC GCAGTCGCGA GCCCTGAAT AAAGCCATAG ATAAATAAAA	8580

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TCCTACAAGC AAGAAAGCTA GTAACTGAAT CTCTCCTGTC AAGAAAGAAA TGATAATAAG	8760
AAAAATATAT AAGGCTGGTA AGACATATTG GTGTAATTGG AATAAAATTC GAAAACCTTG	8820
TTCCAAATTA GCCTGACGCT CCCCTTCATC ATAAGAATTAT ATATAGTTCA AGACATCCTT	8880
TGGTGTAGCG AAAAATTCCA AATCAAACGT ACGAACAAATC GCAATGGTT TAAAAAGAGA	8940
TTTTTGAGCG ACTAAGAATA CCACAAAGAG TAAGAAAGAA AGGAAAAATG TTTGAGGGTT	9000
TGTATGCAAT ATAATCACCT CACTTAATGA AATAAAATAA GCCAATGGAA TCGCTACACC	9060
TGTAATATTA AAAGCAATGG TTCCAAACTC AAGATTCCGA TACATTTGCA CATAATAGGT	9120
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TAAGAAAATTG AAAGTCAAA ACATACTAAT GAAAACCTATC AGTAAACAAA TAGCTGATAT	9240
CCATGGCATE AAGGCTTTA CATCTAAAAT AATTCGTGG GATTGACAC GTGCCTTAAA	9300
CATCCCTACA AACATGCCA AGAACCCCCC AAGACAATAG ACATAAAAAA TAACAATCTA	9360
CGTTCTTTT TCATATTCA TCTCCTTTT CACTTGCTAG ATTTTTGGAT TTCTTTCAA	9420
TCCATTCAAT TACTGGGATG AGAGCAAAGT AGACCCAAAC AAATTGGTCC CTTTGATAGG	9480
GATTAAACCA GCTTAGGTCC ATCCCAATCA GTAGAAATAC GCTGACTAAT AAAGCTATGA	9540
CCACTACATA ATAAATCACT TTATACCTGT TCATCACTCC TCCTCCTCCA AACGAAATAC	9600
CGATTGCACT GTTTCGTTGA AAATTGAGA TATTTTCAGG GCAATGATAA TGGATGGGT	9660
GTACTCATCC CGTTCTAGTA GGCTAATGGT CTGTCGGAA ACCCCTGCCA GTTGGCTAG	9720
GTCGGTTTGA TTGAGACCAT CGCGAGCTCG AAGCTCTTT AGACGATTTT TTAGTTGCAT	9780
GTTACACACC TACTCTCCGT CAAATTCAAC GGTTTGGATA TCCTCAATAC GTTGCAACTT	9840
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CATTCTCTGT AATAATCTCG ACATTTCTCC GTTTCCTTTT TCTTTTCGCT CAAAGCTTTT	10020
GATTTTATTC TCTAGTTCT TGATTTTTT AGAATTATTA GAATAAAAGA AAATCATAAA	10080
TAGTATAAT CCTAGTACCC ACATTATAAC TCCTTTCTGC TTCCTATTTC TTAACCTGAA	10140
TTCATGTAA CATATCTTT TCTTTTGAC AAGTATAGTT GTCAAAAAAA TTATGATTTT	10200
TGTCATTTG CAAAAGAAAA AGTCAGGAG TAGGTTCCCTG ACCACTTTAT CTATCATTAA	10260
TACTCTTCTA AAATCTCTTC AAAACCACGTC AGCTTCACCT TGCCGTAGGT ATGGTTACTG	10320

346

ACTTCGTCAG	TTTCATCTAC	AACCTCAAAA	CCATGTTTG	AGCTGACTTC	GTCAGTTCTA	10380
TCCACAACCT	CAAAACCATG	TTTGAGCTG	ACTTCGTCAG	TTCTATCCAC	AACCTCAAAA	10440
CCATGTTTG	AGCTGACTTC	GTCAGTTCTA	TCCACAACCT	CAAAACAGTG	TTTGAGCAA	10500
CCTGGGCTA	GCTTCCTAGT	TTGCTCTTG	ATTTTATTG	AGTATAAAAT	CCTAGTTTT	10560
CAAAGATTTC	TGAGAAGTTT	TGGCTGATTG	TCTCAAGTGA	CACTTGCACT	TCTTCTCGGG	10620
TTTGGTTGTT	CTTGACCGTC	ACTTGCCGC	TTTCGACTTC	GCTCTCTCCT	AGGGTGATGA	10680
GGGTCTTAGC	CGCAAAGACA	TCGGCTGACT	TGAAGTGAGC	TTTTAGTTA	CGGTTGAGGT	10740
AATCACGCTC	TGCTTGAAA	CCTGTGTTGGC	GAAGAGCCTG	TACCAATTCC	AAGGCCCTGA	10800
TATTTGCCCC	TTCGCCCAAG	ACTGCGATAT	AGACATCTAG	GGCGTTTCG	ATAGGGAGGG	10860
TCACACCTTG	CTTTCAAGG	ATGAGAAGCA	GGCGCTCTAC	ACCAAGTCCA	AAACCAAATC	10920
CAGCAGTTTC	AGGGCCTCCA	AAAGTAAGCAA	CCAAACCATC	GTAGCGACCA	CCCGCACAGA	10980
CGGTCAAGTC	ATTGCCCTCA	ATCTCTGTGA	AAAAGTCGAA	AATGGTGTGG	TTGTAGTAGT	11040
CCAGACCACG	CACCATATTG	GTATCGATGA	TGTAATCTAC	TCCAAGATT	TCCAACATCT	11100
GACGCACAGC	ATCAAAATGA	GCTTGGCTTT	CTTCATCAAG	AAAGTCCAAG	ATAGACGGCG	11160
CATTCTCTAC	TGCCACCTTG	TCTTCTTTTT	CCTTAGAGTC	CAAGACACGA	AGAGGATT	11220
CCTCCAAGCG	ACGTTGGCTA	TCCTTAGACA	AGGTCTCCTT	GAGCGGTGTC	AAATAGTCAA	11280
TCAAGGCTTG	GCGTAGGCT	GCACGGCTCT	CAGGATTTC	AAGAGTGTG	AGGTGCAATT	11340
TGACACCTTG	AATACCGATT	TCCTTCAAAA	AATGGGCTGC	CATAGCGATT	GTTTCCACAT	11400
CGGTAGCTGG	ATTGCTAGAG	CCAAACACT	CAACACCAAT	CTGGTGGAA	TGGCGCAAGC	11460
GCCCTGCCCTG	TGGACGCTCA	TAACGGAACA	TAGGTCCCCT	GTAGTAGAAC	TTGCTTGGCT	11520
TTTGCACTTC	TGGGGCGAAA	AGTTTATT	CCACATAGGA	ACGGACAACG	GGTGCAGTTC	11580
CTTCTGGACG	GAGGTAATA	TGACGGTCAC	CCTGTCTATA	AAAATCGTAC	ATTTCTTGG	11640
TTACGATATC	CGTTGTATCT	CCGACAGAGC	GACTGATAAC	CTCGTAATGC	TCAAAATAG	11700
GCGTGCAC	TTCTGCATAG	TTGAGCGTT	TGAAAATCTC	ACGGCAAAG	CCCTCAACGT	11760
ACTGCCACTT	AGCAGACTCA	GCAGGTAAAA	TATCCTGCGT	TCCTTTGGT	TTTTGTAATT	11820
TCATAGGGAA	TCCTCTTAA	ACTTAATAGT	CTTATTTAC	CATAAAATAGA	GGGATTTAAA	11880
CAGTAAGAAA	AAAATTAGGA	TTTAGATATC	ATTTTGAGA	TTAAGAATTG	TCAAAAAAT	11940
AGCTAGCAAG	GAAAGACCAA	CAAATAGCAT	CCAAGTCAAC	TGTATATTCC	ATACGGCTAC	12000
TAGTGAAAAA	CAAGCTGTT	CCACAGGTAT	GGATAAGGTA	AACAATAGAC	CTAAAAAATT	12060
ACTAGTACGA	GCTAGAACCT	CTGGAGCTAG	ATTTTCATG	AGCATGGCAC	TAATCTTGG	12120

347

TTGAACTTTA CCAGACACAT ACAGAGTAAA GAAGAGAAAT AGCAAACCAA GCACGACTTG	12180
ATTGAATAAA TTAGCCAAAC CAACTAGACT AAGTCCTACG GTCTCCCACA TCATCAATCT	12240
AGGCAAGGAC TGCTTCCCAA AATAATCATT GCCCGTAAGG CTACTGATGA TGACTGATAC	12300
TAAAACACAG AATTGATTGA TAAATAGTGC CTCTGTATAA GAAAATTCA AGAGAGAATG	12360
GCTCAAAAG AAGATATTAT AAATTCCACC CAAAGCGCCA CCCAAGGAAT TAATAAGCAA	12420
GACAGCAAAG AGCATAAAAC CAAAGTTTT CTGTCCACTT TTAAGAAAAA CGAGACGTA	12480
ATTTCCGTAA ATTGTTAGGA ACTGGTCTTT GATAGAAAGC TTCTCATTTC TTAAGTTTC	12540
ACCATCAGCA GATGACATTG ACAGGCTAA TTTGCTTTT CCTAAAAAGA GGATAGTGGC	12600
TGATACTAGG AAAAAGCAGG CATTGATTCC CGCAACGAGA GAAAAATTGT TGACCGATAG	12660
AGCTAAGAGC CAGACTCCGA AACCTTGACC ACCAATAGCT GAAATATAGG TGATGAAC	12720
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ACGCAGGCCA CCTGCAAAT CACTGATGAT ATCACTAATG ACATTGATCA AACACAGGCT	12840
AGAAAAAGCA AAGAGACTAG CTTGCTGAAC AACTAGGGCT GCTAGAAAAA ATAGAACCGC	12900
CTGAAACAAA CCGCTATAGA CCATCCATTG GACCTTGTCC CTCGTGTAAT CTGCCGAAT	12960
CCCTGCAAAA ACTGTAAAGA GGTCGGAAG AATCATGACA ATATTGCCA TAGAACAGC	13020
AAAAGATGCT TGTGACAAGG TCGATGCATA GACGATAAAG ACCAGGTTGA AAATCGAAAC	13080
ACCAAAAGCA TTGACACACC GTGG	13104

(2) INFORMATION FOR SEQ ID NO: 35:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 19250 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 35:

CGGGGCAAAT AGTTTGAACTTTCATCAT TTTCTCCTTT AAAACTTTCT CTCCATTATA	60
GAECTTTTC AGAAAGTTGT CAACAGAATT TTCAGAATT TTGAAAATTAA TTTTCAAAAC	120
AACATCTTG CAAAAAATAT GAATATCGTA AGCGCGTCAT AACAAAGGTAT CTATCATTCA	180
TGGAGCTCCT CCTGTATACT ATTAGTAAAG TAAATATTGG AGGATATTTC AATGCCACAA	240
CCTATTGTTCTGCTGAGAT TCCACAATCT CGTCGTTTG ATTCTAAAAA GAGAAATGAT	300
ATTCTCTTA AAATTCGTAT TGGCAAGCTT GAAAGTAAGTT TTTTCAATC TCTCAATCTC	360

348

GAAATGATAG AACAGCTTTT GGATAAGGTG TTGCTCTATG ACAATTCACTC TATCTAGCCT	420
AGGGCAGGTC TATCTCGTGT GTGGGAAAC TGATATGAGA CAAGGAATCG ATTCACTGGC	480
TTATCTCGTT AAAACCCACT TTGAAATTGGA TCCCTTCTCC GGTCAAATCT TTCTCTTTG	540
TGGTGGACGT AAAGACCGCT TAAAGTCCT TTACTGGGAT GGTCAAGGAT TTTGGCTACT	600
ATATAAACGC TTTGAGAACG GCAGACTGAC TTGGCCCAGT ACAGAAAAGG ATGTCAAAGC	660
TCTCGCACCT GAACAAGTAG ATTGGCTGAT GAAAGGCTTT TCTATCACTC CAAAAATATA	720
GTAGATTGAA ACTAGAATAG TACACCTCTG CTTCTAAAAC ATTGTTAGAA ATCGATTTA	780
CTGTCCTGAT CGATTGTCC TGGTATTATT TCATTTACT ATAAATCCAT CAGAAAGTCG	840
TGATTCTAT TGAAATGAGG ACTTTCTTT TATACTCATC TGCTTCAAA AAGCACTCTA	900
GTCCATCTCC GATTAACCGT GGACTTATC ACCTCCTTCT CCAGTCCTTG TATAACATCT	960
TGAAGTTGAT TCATGACATC TCCAAAGTT CGAAAGGCTT TATTCTAAA TCCACGTTA	1020
CGAACATCTT TCCACACTTG TTCAATGGGG TTCATCTCTG GTGTGTATGG AGGAATAAT	1080
GCAAAGCCAA TATTAGTCGG AATCTTAAG GTACTTGATT TATGCCATAT AGCATTGTCC	1140
ATAACGACTA AAAGATAATC ATCTGGATAA GCTTGTGAAA GCTCCTATTG CTAAAGCCCC	1200
TTTATAACCT CTTGGCAGAG AGACTATTGA CTCAGCCCTT ACTTCATGCG GATGAAACCT	1260
CCTATCGGGT TCTAGAGAGT GATAGCCATC TGACCTACTA TTGGACTTTT TTGTCAGGTA	1320
AAGCAGAGAA ACAAGGGATT ACGCTTAC ACCATGATCA GTGTCGAAGT GGTCAGTAG	1380
TACAAGAATT CCTAGGAGAT TATTCTGGCT ATGTTCATTG TGATATGTTG CGGCAGTAAC	1440
TTAGGACTTT AGTCCTCTAG TTCTGCCTAT GCGATAGCAG TCCAAGGTTT AGGAGTAAGG	1500
CGACGCTAAG CTTGGTAAAC TCGAACAGC TAGAAGCTTA TCGTCAACTG GAAGAAGCTG	1560
CACTTGTGG ATGTTGGCG CATGTGAGAA GGAAGTTTT TGAAGTGCC CCCAAGCAAG	1620
CAGATAAACATC ATCCCTAGGA GCTAAAGGTT TAGCCTATTG TGATCAGTTA TTTTCCTGG	1680
AAAGAGACTG GGAGGCTTTG CCAGCTGATG AACGGCTACA GAAACGTCAA GAACATCTCC	1740
AACCCCTACT GGAAGACTTC TTTGCTTGGT GCCGTCGTCA GTCAAGTTTA TCGGGTTCAA	1800
AACTAGGAAG GGCAATTGAA TACAGCCTCA AGTATGAAGA AACCTTAAG ACCATTTAA	1860
AAGACGGACA TCTGGCTTT TCCAATAATC TAGCTGAACG CGGCATTAAA TCATTGGTTA	1920
TGGGACGGAG TAAAAGAGTC CACTGGACTC TTTAGCCTA AGCTCAGTTT AAAAAAACGA	1980
GGGTGGTTAT TTTAAAAAA CGCAGGGTGG TTATTTCTC AAAGTTTGAG AGGAGCTAAA	2040
GCAAGAGCTA TTATTATGAG TTGTTGGAA ACAGCTAAC GTCATCAATT ATAGTGCCTT	2100
GAATCTATAA CAGTACGCAT CGACTGCTAA AATATTCTA TAAATCAATT TTCTTTCTC	2160

AATCGATTG TTCATATCTT ATTACAATCC ATTATAAATA GCGAGAAATA TCTATCCTAT	2220
CTTCTAGAAT GTCTTCCAAA CGAGGAAACT CTCGTAAACA AAGAGGTTT AGAGGCCTAT	2280
TTACCGTGG A CTAAAGTTGT ACAAGAAAAG TSCAAATAAG AAATCTCCAG ATTAGGAAC	2340
ATATATGAGT TCTCTAGTCT GGAGATTTT CAATAGACTT CGTTATTGGG CGGTTACTTT	2400
CGAAACTTTG AAAACTTCAA AAAACGGATT TTATCGCTC TGAACATCAA AAAAGAAAGG	2460
ACGAAATTG TCCTTCTCA AGCTTAGCTT TTCTCAACC CACTACAGTT GACAAAGAGC	2520
CTTTTATTCT ATCAAACATG AAGCGAAAA ACAAGCCAA AATCCGATAG AATGGCTATC	2580
CCTCGACTAT CAAGTAAGAC ATTTCCATCA AATACGTTCA ATTTTACTCT TGTTCTACTA	2640
AGAATTAATC ATCTCGTTT GATTTATTAA AAATATACAA TTCAGCTTT CCTCCAAACT	2700
ATTTTATCCA CTATCCCTGT ATAGCTCTGT ATTATCTTAA CAACTTTAGT AGAGACATT	2760
TCCTCAACAT AATCCGGAAC CGGTAAATCCA AAATCCTCAT CTTGTGCCAA GCTAACAGCA	2820
GTTTCAACTG CTTGAAGAAG AGAATTTCA TCAATGCCTG CAAAATAAA TCCTGCCTTA	2880
TCTAAGGACT CAGGACGTTC TGTACTTGTA CGAACATATA CAGCGGGAAA AGGATAACCT	2940
TGACTAGTAA AGAAACTACT TTCTTCCGGT AAAGTCCCG AATCAGATAC TACAACAAAT	3000
GCATTCTCATCT GTAAACAATT ATAGTCATGG AATCCTAGTG GCTCATGCTG AATCACACGT	3060
TTATCTAGTT TAAAACCGCT CTCTTGAGC CTTTTCTTTG ATCTAGGATG GCAAGAATAT	3120
AAGATTGGCA TATTATACCTT TTCACTAAT TGATTAATTG CTGTAAAGAG AGAAATAAAA	3180
TTTTTATCTG TATCAATATT TTCCCTCACGG TGAGCTGAAA GTAAGATATA ACCTCCTTT	3240
TTCAATCCC AACGTTCATG GATATCTGAA GACTCAATAG CAGATAAATT TTTATGTAAC	3300
ACTTCTGCCA TAGGAGAACC AGTTACATAT GTGCGCTTT TAGGTAACACC ACACATGT	3360
AAATACTTAC GTGCATGTTC AGAGTATGCT AAGTTAACAT CTGAAATAAC ATCAACAATC	3420
CGACGATTAG TCTCTCCGG TAGGCACTCA TCTTACAGC GATTGCCAGC CTCCATATGA	3480
AAAATTGGAA TATGTAACG CTTGGCAGCA ATAGCTGATA AACAGAAATT TGTATCCCT	3540
AAAATCAATA AAGCATCTGG TTTAATTGAT TTCACTCAATT TGTATGAAGT ATTAATAATA	3600
TTCCCTACAG TAGCACCAAG ATCATCTCCA ACAGCATCCA TGTATACGTC CGGAGTGTCT	3660
AACCCCTAAAT TATCAAAGAA AATACCATT AAATTGTAAT CATACTTTG TCCAGTATGT	3720
GCCAAAATAA CATCAAATAA CTTTCGACAT TTAGTGATAA CACTACTAG ACGTATAATC	3780
TCTGGACGTG TTCCCACAAT AATCAATAAC TTAAGTTGC CATTATCTTT AAAGTGAATA	3840
TCACTATAAT CTGTCTTAAT TTTCATTTAT TTCTCCACTT GTTCAAAAAA AGTATCTGGA	3900